REMOVAL OF SOLUBLE BOD 5 IN PRIMARY CLARIFIERS

bу

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FOREWORD

When energy and material resources are extracted, processed, converted, and used, the related pollutional impacts on our environment and even on our health often require that new and increasingly more efficient pollution control methods be used. The Industrial Environmental Research Laboratory - Cincinnati (IERL-Ci) assists in developing and demonstrating new and improved methodologies that will meet these needs both efficiently and economically.

Removal of soluble BOD5 in primary clarifiers presents research findings related to means and methods for improving the efficiency of primary sedimentation systems treating pulp and paper wastes. The possibility of greater BOD removal was investigated by examining effects of flocculants and coagulants in removing soluble and colloidal constituents of such wastes. The report also discloses that certain process waste streams may be antagonistic to efficient sedimentation and should be treated separately from total mill wastes. This information will be of interest to segments of the industry wishing to upgrade primary treatment efficiency to reduce loadings on secondary treatment. The findings are also applicable to treatment of waste streams not containing significant additives, as the tissue industry, wherein effluent guide-line allowances may possibly be satisfied by efficient primary treatment. For further information contact the Food and Wood Products Branch, Industrial Environmental Research Laboratory-Cincinnati.

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ABSTRACT

Primary clarifiers used in the pulp and paper industry show wide variations in BOD removal. This project was directed to evaluating means for increasing BOD removal and developing an understanding of the mechanism. Such understanding should permit optimization of BOD removal during primary clarification.

A survey involved twelve mills, nine with primary clarifiers, to obtain data on total and soluble BOD₅, COD, suspended solids, and color. These data were used to select mill effluents for additional study.

Laboratory studies showed that, with the proper flocculating agent and operating conditions, the BOD₅ concentration of some mill effluents could be markedly reduced during clarification. These findings were confirmed by pilot-scale clarifier field trials at two mills.

Measurement of sludge volumes and sludge dewatering characteristics resulting from chemical treatment were outside the objectives of this study.

Gel chromatography studies showed that considerable low molecular weight biodegradable residues along with colloidal material were also flocculated and removed. Studies with model compounds indicated that increased removal is apparently related to the pH of the solution and to the functional groups (hydroxyl, phenolic, and carboxylic), chain length, branching, and solubility of the compound.

Chemical costs may range from 3¢ to 10¢ per 1000 gallons of feed to the clarifier. Decreased cost and increased operating efficiency in total treatment systems, as a result of improved clarification, could more than justify this chemical charge. Minimum levels of chemical costs will depend upon minimizing or eliminating overflows or spills of colloids and dispersants detrimental to clarifier efficiency.

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Ralph H. Scott, Chief of Pulp, Paper and Wood Staff, Food and Wood Products Branch, IERL-Ci, EPA, as Project Officer, has had responsibilities for the immediate supervision of this project in accordance with the objectives of the program.

SECTION I

INTRODUCTION

The overall degree of biochemical oxygen demand (BOD) removal actually observed in most conventional out-plant clarifiers treating total mill effluents has been relatively small (5-15 percent) and has seldom extended to levels that would significantly improve effluent quality. A search of the literature disclosed few published investigations of the mechanisms for the BOD removal in clarifiers. Only two (1,2) gave data indicating possible sources of BOD that is removed during clarification. The State of the Art Review of Pulp and Paper Waste Treatment by Gehm (1) attributes the BOD to "the high content of suspended organics and small quantities of dissolved organic matter" in certain types of waste and to the "fact that the oxygen uptake rate of fiber is slower than that of dissolved materials, since it must first be liquefied before oxidation can take place." Das and Lomas (2)point to glucuronoxylans and araboglucuroxylans as water soluble hemicelluloses (from the beating of pulp) that reduce the BOD when removed with the floc. One of the first goals of Phase I of this research program was, therefore, to evaluate by a controlled analytical study the removal performance for total and "soluble" BOD5 in different clarifiers at pulp and paper mills. Surveys of mill waste flows and clarifier effluents were conducted at 12 mills owned by 11 companies. The analytical data developed during this survey established the basic program of the investigation and the definitions of the terms "soluble" and "total 5-day biochemical oxygen demand (BOD₅)" to be used throughout the balance of this study.

Three different classes of BOD-forming components in the various waste flows were identified:

- 1. Substantial quantities of the BOD being removed in clarifiers are obviously suspended fibers and related solids in various stages of physical, chemical, and biological degradation. These suspended particles may originally have low levels of solubility and low BOD but can undergo degradation to produce soluble substances responsible for BOD. This fraction comprises the principal source of BOD readily removed in conventional clarifiers.
- 2. Advanced degradation of cellulose and hemicellulose particles and the presence of papermaking additives such as starch may provide a substantial amount of soluble, high molecular weight components with biochemical demand for oxygen. Most of these kinds of substances can be surface-adsorbed and removed by advanced clarification procedures. The further development and field testing of routes to removal of this fraction comprised a principal area for development under Phase II of this project.

3. A third fraction of the BOD contained in pulp and paper waste waters is comprised of low molecular weight degradation products of carbohydrates, such as acetic acid, aldehydes and alcohols, which are not readily amenable to physical adsorption and flocculation. Removal of these components has not been reported for primary treatment if we except those cases of prolonged holding time or other conditions where advanced stages of microbiological action are evident in the clarifier.

The fundamental physicochemical mechanisms of flocculation of pulp mill white water fines studied by Williams ($\underline{3}$) and the work by Back ($\underline{4}$) on the nature of wood components dissolved in the pressure refining of mechanical pulps provide information on a possible approach to the development of an understanding of the mechanisms involved in the removal of BOD components from waste streams. Their ideas were used to develop the jar testing procedures used in our laboratory for the optimization of the removal of BOD5 constituents from various total mill effluents.

Phase I studies were funded jointly by a group of U.S. pulp and paper companies (Appendix A), the U.S. Environmental Protection Agency (EPA) (Grant No. 803 119), and the Wisconsin Department of Natural Resources (Ref. No. 8100).

For a supplementary and confirming Phase II program, small-scale pilot equipment was used for on-site comparative studies, paralleling commercial clarifiers operating at two mills representative of a broad segment of the pulp and paper industry. Additional funds were supplied by the U.S. Environmental Protection Agency under an extension of the above grant and by The Institute of Paper Chemistry. This confirming program of study was designed to evaluate the operating parameters developed in the earlier Phase I laboratory research under mill conditions. The first trial was conducted at an integrated kraft mill and the second at an integrated mill producing pulp by a recently developed chemimechanical (CM) pulping process.

Laboratory jar tests were also used to evaluate the flocculation and clarification characteristics of individual sewer discharges at these two mills. We studied the effects of processing each of these streams separately and we could also check the effect each had on the overall BOD_5 efficiency of the clarifier when processed as a part of the total mill effluent. Additionally, the jar tests were used to test the removal of soluble model compounds (acetic acid, aromatic dicarboxylic acids, etc.) from solution with the various flocculating agents in order to help elucidate the mechanism of BOD_5 reduction.

An economic evaluation of the data of two field trials and the results of the jar tests conducted on individual streams show the savings which might be realized if the relatively small volumes of stronger pulping effluents, containing lignin compounds with dispersant properties, were not treated in the clarifier.

SECTION II

SUMMARY AND CONCLUSIONS

- l. A large portion of the BOD entering the primary clarifiers processing pulp and paper manufacturing waste waters are solubles that can pass through a $0.45~\mu m$ filter.
- 2. These components are generated from wood in pulp and papermaking steps, including wood handling. They may also come from papermaking additives such as starch or sizes. These materials can be degraded within the mill or during out-plant treatment.
- 3. Laboratory jar tests with various chemical flocculating agents and operating parameters showed that significant quantities, ranging to 20 percent or more, of the soluble BOD₅ and color could be removed from wastes from mills surveyed in Phase I studies. These findings were confirmed by pilot scale field trials at Mosinee Paper Company at Mosinee, Wisconsin, and at the Combined Locks mill of the NCR Corporation in Combined Locks, Wisconsin.
- 4. Jar tests with the individual waste streams making up the total mill discharge have shown that certain streams contribute both high BOD_5 levels and chemicals that are detrimental to the efficient operation of the clarifiers. Elimination of these streams from the out-plant clarifiers could provide increased quality to the mill discharge. It would also decrease the load on the secondary treatment plant. The streams that were removed from the influent to the clarifier could be treated separately in small clarifiers or by reverse osmosis, ultrafiltration, or other small unit processes.
- 5. The mechanism for the removal of soluble BOD_5 is very complex due to the heterogeneity of the wastes treated in the clarifiers. There was no apparent relationship between the initial soluble BOD_5 concentration and the ultimate removal of the soluble BOD_5 components by the clarification process.
- 6. Gel chromatography analysis of effluents before and after coagulation with iron and pulp fines showed a significant number of low molecular weight compounds were removed. Model compound studies showed that many organic acids including some quite soluble could be partially coagulated in an iron-pulp fines system as well as substances which do not form salts with iron.
- 7. Evaluation of the benefits and costs of achieving improved performance of clarifiers requires detailed engineering studies to fit individual mill situations. Such studies were beyond the scope and objectives of this study. However, it was apparent that total chemical costs, including the chemical additives which may be used in existing practices at these mills,

might range from 3ϕ to 10ϕ per 1000 gallons treated. Efficient use of these chemicals would be balanced by benefits and cost reductions in the overall systems of waste treatment.

8. Good clarifier performance and low chemical costs could only result from adequate control of mill overflows and spills so that strong colloids or dispersants do not enter the clarifier.

SECTION III

RECOMMENDATIONS

These studies have shown that with the proper selection of flocculating agents and operating conditions, soluble BQD₅ concentrations in waste streams can be markedly reduced (10-50 percent) in the primary clarifier.

Collapse of the double layer surrounding the charged molecule, bridging, and decreased solubility after treatment with the flocculating agent are mechanisms which could be expected to contribute to the removal of both total and soluble BOD₅ components, as well as color, from the effluents.

- l. We need to know more about the constituents and characteristics of different kinds of sludges those from high and low levels of soluble BOD_5 removal and their effect on the efficiency of clarification in mills processing under different conditions. This could add considerably to our knowledge of the mechanisms involved. The influence of individual mill waste streams in clarifier sludge performance also needs further study.
- 2. Many of the individual sewer discharges were found to have adverse effects on the efficiency of the clarifiers and also to have a toxic effect on the 5-day BOD test. Removal of these streams from the clarifier inflow could markedly increase the efficiency of both the primary and secondary (biological-type) treatment systems. Diversion of these flows, which are usually of small volume, to other treatment systems is recommended.
- 3. Selection of the streams to be "eliminated" could be undertaken under laboratory conditions with a combination of jar and BODs tests. Processes for treating the eliminated streams by separate in-plant clarification, reverse osmosis, ultrafiltration, or the training of plant personnel to reduce the discharge of the detrimental stream would follow the identification of these major sources of difficulty.

SECTION IV

MILL SURVEYS

EQUIPMENT AND PROCEDURES

Weekly samples from mills using clarifiers consisted of two gallons of the influent to the clarifier and a single gallon of the clarifier effluent. Where possible, both samples were collected as 24-hour composites. Mills not using clarifiers sent in two gallons of the effluent they were discharging into ponds or aeration lagoons. Table 1 describes the types of mills and their facilities for pulping, bleaching, and effluent treatment; this table is based on data supplied by the mill staff.

As soon as the samples were received in the Institute laboratories at Appleton, Wisconsin, the influent samples were composited, well mixed and immediately transferred to one-gallon plastic bottles. One bottle was filled to the brim, set aside to settle for one hour, and the top 6 inches of supernatant was withdrawn using a light vacuum and a short U-bend at the end of a glass tube. This permitted removal of the top layer of "clear" fluid without drawing the "settled" material from the bottom layer. A second bottle of the influent and also a bottle of clarifier effluent were separately mixed and analyzed "as received."

Samples of the "as received" and "settled" material were pressure filtered through various porosity filters. These filters were 1.2- μ m and 0.45- μ m pore size Metrical membranes from Gelman Instruments of Ann Arbor, Michigan and a 0.10- μ m plastic filter from Nuclepore of Pleasant, California.

Suspended solids, BOD₅, and chemical oxygen demand (COD) were determined on the initial samples and on all fractions.

Samples for suspended solids were processed on 1.2- and 0.45- μ m filters on a Gelman pressure funnel at 6.9 kg/cm² (100 psig) with N₂ and dried at 103-105°C (218-221°F) for one hour. For biochemical oxygen demand we used the standard APHA Method (5), keeping the samples at 20°C (69°F) in a water bath and determining the dissolved oxygen with a Weston and Stack probe. The COD was determined by the dichromate oxidation method of APHA (6). Color was determined at pH 7.6 using the National Council for Air and Stream Improvement method (20).

Some waste samples filtered slowly through 0.45- and 0.10-µm filters. To avoid filter clogging, a pad of filter aid (Celite, AR, Johns-Manville) was used on 1.2-, 0.45- and 0.10-µm filters for these wastes. The filter-aid pad was formed from a 1-percent slurry of Celite AR poured onto the filters and then dewatered. Analysis of these samples with and without the use of the filter aid (Table 2) indicated that the filter aid did not adversely affect the analytical accuracy. Therefore, in later work, the filter aid was

TABLE 1. DESCRIPTION OF MILLS SURVEYED

Mill	Type of cook	Tons/ day	Bleaching sequence	Effluent sewers	Volume, mgd ^a	Clarifier additives
Ą	Ammonium bisulfite, aspen	NA	NA b	Pulp mill wash water	2.6	Gendriv 162 2 mg/1
щ	Acid sulfite	NA	NA	NA		No clarifier
U	Groundwood	225		Coater Paper machines Boiler room & treatment plant	0.19 2.3 0.2	Cationic polymer 1.7 mg/l
Q	Acid sulfite (pH 2.5) 66% softwood 34% hardwood	115	Single stage calcium hypo- chlorite	Evaporator condensate Decker Bleach washer Paper mill Miscellaneous	0.1 1.0 1.0 1.0 1.0	None
EE EE	Kraft, groundwood	NA				No clarifiers
ͱι	Chemimechanical, also NSSC and kraft fibers used in papermaking	350	High density peroxide	Wet room CM No. 6 Main No. 5 Old filter backwash New filter backwash	2.16 1.11 1.117 1.10 0.53 0.40 0.43	None
t)	None-deinking mill	150	С-ЕН-Н at volume ratio 90:5:5	Decker deink Bookstock deink No. 3 & 4 paper machines No. 5 Paper machine Beater room additives Chlorination waste water Misc. (water treatment plant sludge, etc.)	0.50 0.10 0.10 0.10 0.10 0.10 0.10 0.10	Anionic polymer 1.5 mg/l

See end of table for footnotes.

TABLE 1 (Continued). DESCRIPTION OF MILLS SURVEYED

Mill	Type of cook	Tons/ day	Bleaching sequence	Effluent sewers	Volume, mgd	Clarifier additives
щ	Kraft Pine with 10% hardwood (oak)	370	None	Paper machine Salvage filter Excess screen room white water Water treatment blowdown Utilities department Kinney strainer reject Hot pond overflow	44 40 mm 1 00 000000000000000000000000000000	None
н	Kraft Pine, Douglas- fir, western larch, spruce	1150	СИНО	Clear water Pulp mill & recovery Bleach plant, paper mill & recaustizing	3.5 5.0 ^c 7.3 ^c , ^d	None
JJ	Converting mill	205	None	Mill effluent Sludge from water treatment	4.75 ^h 0.08 ⁱ	None
وا	Groundwood	455	None	Paper machine stock room and coating plant Paper machine Off-machine coater Groundwood operation Cooling & miscellaneous	w4001 0%%00	Polymer, 0.5 mg/l
×	Kraft Pine	230	None	Paper machine High density Pulp mill Foul evaporator condensate Soda recovery Digester blowdown condensate Digester room 5th Stage brownstock washer	10.0° 1.55 0.0° 0.0° 0.0° 0.0°	Hercofloc 812.3, 0.75 mg/l
0			9			

Defoamers and detergent added.
Alum and silicates added.
Wet strength resin, defoamer, talc, animal glue, slimicides, rewetting agent, dryer release, and organic dispersants added.
Alum and chlorine added. anged = Million gallons per day.

NA = information not available.

Deformers added.

Pitch control chemical added.

Estimate. used as a standard practice with the precaution of first checking the effect on one or two samples of each waste stream. Eventually the exploratory studies with various grades of filters resulted in adoption of the 0.45-µm porosity as a standardized procedure.

TABLE 2. ANALYSIS OF FILTRATES PREPARED WITH AND WITHOUT FILTER AID^a

	BQD ₅ , mg/l				COD, mg/l			
	1.2	2 jimp	0.45 μm ^c		1.2 µm		0.45 μm	
Sample	With	Without	With	Without	With	Without	With	Without
F	190	178	182	176	464	436	409	369
	74	82	70	80	212	216	199	191
Н	32	34	31.	32	133	123	108	114
	48	42	44	46	105	126	118	106
В	770	778	672	675	3741	3650	3636	3615
	1246	1230	1218	1184	5363	5320	5190	4890
C	166	192	188	184	352	362	329	329
	184	181	168	174	316	316	275	292

 $_{h}^{a}$ Celite (AR) BOD₅ = 0.04 mg/g.

A few samples were also processed through a Dia-Flo (Amicon Scientific Systems) stirred ultrafiltration (UF) unit with a nominal 100-A (0.01 μm) pore filter. The results were compared with those from the 0.45- and 0.10- μm filters (Table 3). Since no differences were apparent in either BOD5 or COD, the more time consuming UF procedure was not routinely used.

TABLE 3. COMPARISON OF ULTRAFILTRATES FROM A 100-A MEMBRANE WITH FILTRATES PREPARED WITH FILTER AID OVER 0.45- AND 0.10-µm FILTERS

	BOD, mg/l			
Sample	SUF ^a	S045F	SOLOF	
F	259	310	266	
	272	256	267	
H	30	43	32	
	59	48	50	
JJ	40	38	35	
	35	40	40	

aSUF - Dia-Flo stirred ultrafiltration cell with 100-A membrane.

b1.2- μ m BOD₅ = 1.92 mg/sheet.

 $^{^{\}circ}$ 0.45- μ m BOD₅ = 0.62 mg/sheet.

DATA AND DISCUSSION

In order to establish the various total and soluble components of BOD5, the incoming samples were analyzed "as received," "settled," and after passage through either 1.2-, 0.45-, or 0.10-µm filters (Figure 1). Throughout this section of the report, the designations are as follows:

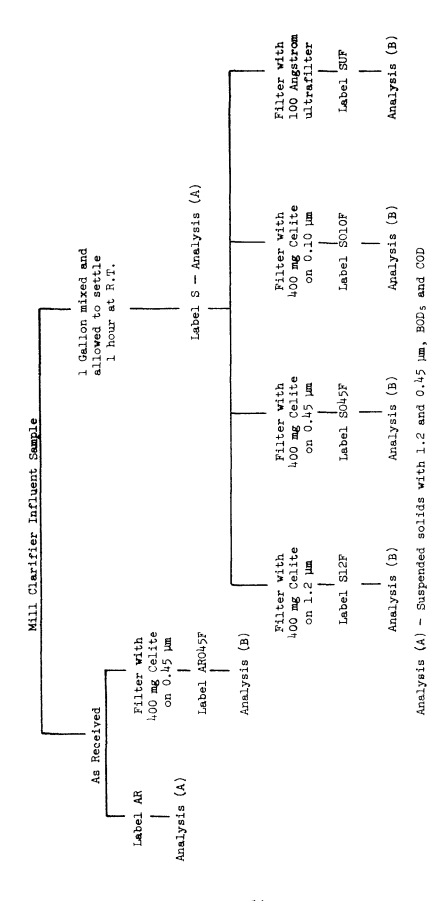
AR ARO45F	As received As received, filtered through 0.45-µm filter with a Celite
	overlay
S	Settled
S 12	Settled and passed through a 1.2-um filter
S12F	Same as S12, except with Celite overlay
SO45	Settled and passed through a 0.45-µm filter
S045F	Same as SO45, except with Celite overlay
SOLOF	Settled and passed through a 0.01-um filter with a Celite
	overlay
SUF	Settled sample filtered through a 100-A UF unit

Data in Table 4 are averages of the six samples from each mill, along with high and low values (range) for each set. The values in parentheses are the percentages of the constituent in the sample <u>after</u> treatment. BOD₅ decreases as filter pore size decreases (Figure 2). The data in Table 4 and Figure 2 can be summarized as follows:

- 1. Comparing "as received" and "settled" values, we find a marked reduction in BOD₅ (to 62 percent) after simple settling (JJ).
- 2. The 0.45-µm filter was, for all practical purposes, the best filter for the removal of "insoluble" BOD₅. This filter was used for further studies of "soluble" versus "total" BOD₅.
- 3. Seven percent additional BOD₅ could be removed from the JJ mill samples by passing them through the 0.45-µm filter.
- 4. The COD values paralleled, but did not directly correlate with, the BOD₅ values.

In order to evaluate the operation of the mill clarifiers we analyzed clarifier influent and effluent samples in the same manner. Data for these samples "as received," and filtered through 0.45- μ m filters, are given in Table 5 (BOD), Table 6 (COD) and Table 7 (suspended solids). In order to simplify the presentation, the "as received" values are plotted in the same sequence in Figures 3, 4, and 5. Marked reduction in total BOD₅ was achieved in clarifier JJ. All clarifiers except those of Mills I and J achieved 50-90 percent removal of suspended solids.

Unfortunately, the values for constituents and removals are quite erratic, probably due to a time lag for arrival of these constituents at the sampling points. Although most samples were 24-hour composites, compensation for the effect of the clarifier "holding time," in some cases on the order of 18 hours and in most cases at least 4 hours, was not built into the sampling system. While this would have little effect on the monthly mill averages, it had a marked effect on the samples sent to our laboratories (Table 8). We found major differences in pH, BOD₅, and COD for the two streams in spite of



Schematic for processing clarifier samples for soluble BODs study. Figure 1.

Analysis (B) - BODs and COD

Suspended solids, mg/l ARO45 S12 1680 556-2333 182 124-241 137 108-206 136 36-272 110 25-245 66 36-108 4165 2473-6086 1218 216-2332 1126 752**-**1780 1242 888-1848 853 360**-**1628 690 290-1118 441 262-548 412 232**-**872 235 154-408 318 192-384 240 182-376 3984 2453-6075 644 270-1034 1092 508-1788 1097 676-1461 633 260**-**1456 TABLE 4. ANALYTICAL DATA FOR TREATED AND UNTREATED SAMPLES DURING SYALUATION OF METHOD FOR "SOLUBLE" BOD: 136-2110 278 138-384 177 97-222 4032(83) 2132-5355 1066(57) 892-1396 1388(65) 491-2495 1250(62) 787-2106 1146(39) 574-1521 260(36) 171-372 248(42) 167-390 395(22) 309-498 331(20) 336(47) 278-373 100(37) 75-118 1207(65) 960-1538 1349(67) 801-2186 1144(54) 588-2350 1201(41) 574-2565 283(40) 188-382 246(42) 175-405 414(23) 302-544 366(22) 191-508 329(46) 108(49) 1218(65) 1000-1529 4370(90) 2127-5765 1360(67) 801-2186 1214(42) 574-1633 1668(78) 597-2975 448(25) 304-598 370(52) 275-586 387(23) 212-527 107(39) 83-130 1262(68) 1016-1624 4548(94) 2457-6000 1821(62) 812-2796 1523(75) 854-2472 1910(90) 914-3195 427(60) 313–621 589(33) 442-696 522(31) 23**8-**762 168(62) 102-246 383(66) 502(70) 357-756 1863 1402-2286 2918 1702-3990 1763 1156–3284 2132 1065-3388 715 411-1138 1684 802-2268 940-3494 712 608-780 272 174-416 583 477-724 162(59) 100-239 409(66) 222-730 412(84) 251-591 188(64) 498(63) 255-687 65(44) 24-132 140(37) 72-220 36(47) 24-50 536(68) 286-711 404(65) 225-728 188(64) 112-234 147(41) 71(48) 28-126 1014(90) 756-1338 172(63) 112-254 538(68) 270-692 451(92) 272-663 191(65) 119-226 154(43) 73(49) 39(52) 27**-**52 1076(96) 726-1407 354(71) 254-510 208(75) 128-273 204(88) 120-263 456(93) 285-665 235(80) 125-281 197(55) 84-393 630(80) 286-847 575(92) 311-765 88(59) 51-144 49(64) 26-70 1081(96) 602-1359 315(64) 215-422 168(61) 108-266 ARO45F 295 156-394 230 153**-**310 149 100-212 276 222-327 4.77 490 3.4-10.2 298-710 359 121**-**613 9.93 787 6.6-11.5 482-927 76 44-110 8.24 7.20 6.16 5.2-6.6 5.43 3.6-7.9 8.74 7.0-8.1 6.98 6.7-7.6 7.46 6.9-8.7 6.68 6.5-6.8 Hd

(% Remaining = settled (or filtered) value x 100)

12

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В

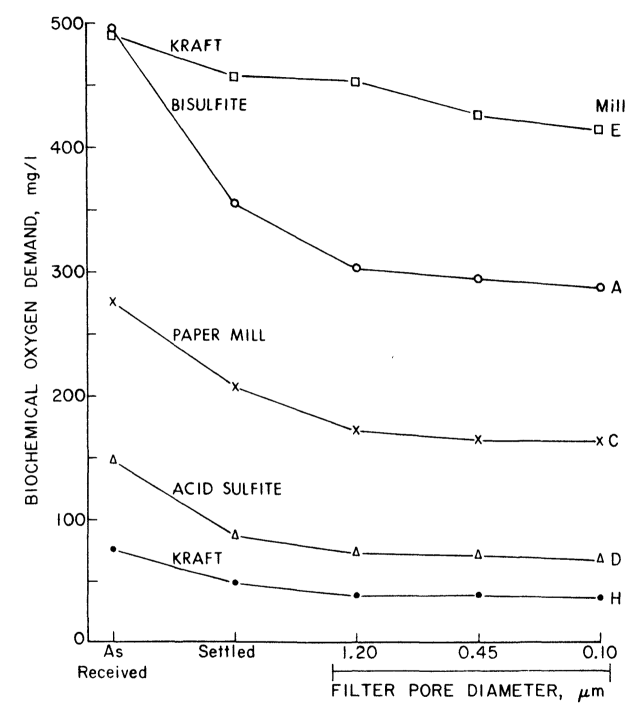


Figure 2. The separation of insoluble from soluble BOD_5 by filters of various pore sizes.

TABLE 5. BOD5 IN CLARIFIER INFLUENT AND EFFLUENT SAMPLES ("AS RECEIVED" AND FILTERED THROUGH A 0.45 μm FILTER)

	As received (total)			Filtered (soluble)				
Mill	Influent, mg/l	Effluent, mg/l	Removal,	Influent, mg/l	Effluent, mg/l	Removal,		
A	495	341	31	303	304	-0.3		
В	No clari:	fier						
С	264	192	27	145	138	5		
D	156	90	42	7 5	59	22		
E	No clari	No clarifier						
EE	No clari:	fier						
F	378	322	15	179	200	-1 2		
G	787	645	18	536	432	45		
H	76	51	33	39	33	15		
I	622	557	10	371	247	34		
J	230	152	33	134	124	8		
JJ	122	37	70	38	32	16		

TABLE 6. COD IN CLARIFIER INFLUENT AND EFFLUENT SAMPLES ("AS RECEIVED" AND FILTERED THROUGH 0.45 µm FILTER)

	As received (total)			Filtered (soluble)		
Mill	Influent, mg/l	Effluent, mg/l	Removal,	Influent, mg/l	Effluent, mg/l	Removal,
A	1863	1328	29	1207	1236	-2
В	No clari:	fier				
C	712	429	40	283	286	-1
D	583	386	34	246	254	- 3
E	No clari:	fier				
EE	No clari:	fier				
F	1695	642	62	408	399	2
G	2918	1606	45	1201	1149	4
H	272	188	31	108	98	9
I	2132	2013	6	1144	1104	14
J	715	433	40	351	284	19
JJ	691	126	82	88	84	5

the fact that both were collected within the same time period. This lack of continuity in the two samples could explain not only the lack of removal observed at times but could also be the reason that our studies showed the higher or lower removal values than did general mill experience.

TABLE 7. SUSPENDED SOLIDS IN CLARIFIER INFLUENT AND EFFLUENT SAMPLES REMOVED

BY A 0.45 µm FILTER

Mill	Influent, mg/l	Effluent, mg/l	Removal,
A	441	63	86
В	No clarif	ier	
C	1126	248	78
D	235	77	67
E	No clarif	ier	
EE	No clarif	ier	
F	1187	213	82
G	4165	1034	75
Н	240	99	59
I	853	587	31
J	690	340	47
JJ	581	50	91
			

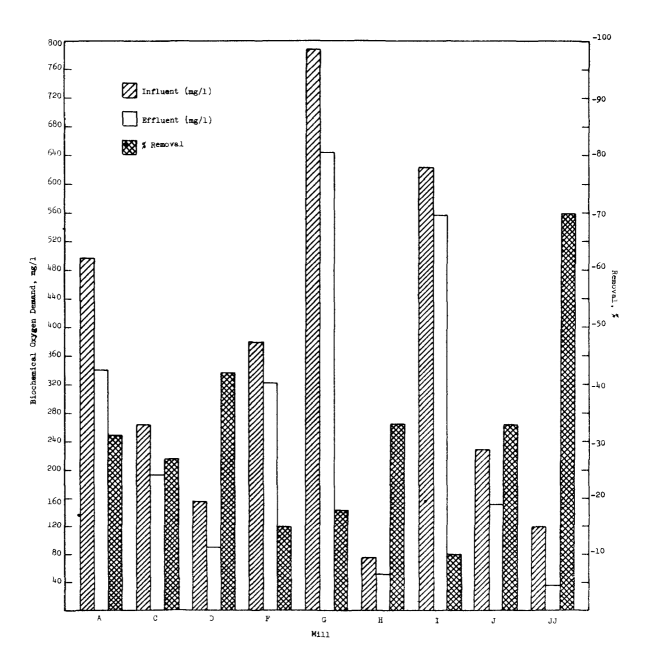


Figure 3. Total biochemical oxygen demand in clarifier influent and effluent samples.

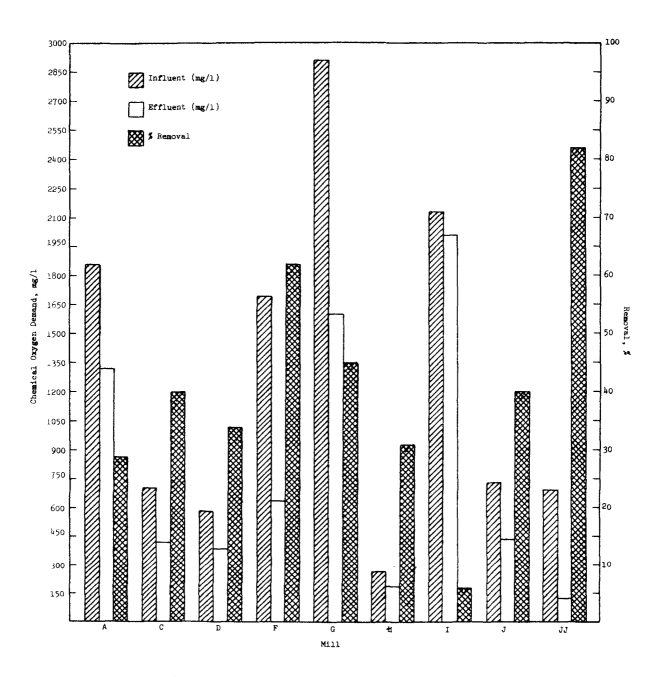


Figure 4. Total chemical oxygen demand in clarifier influent and effluent samples.

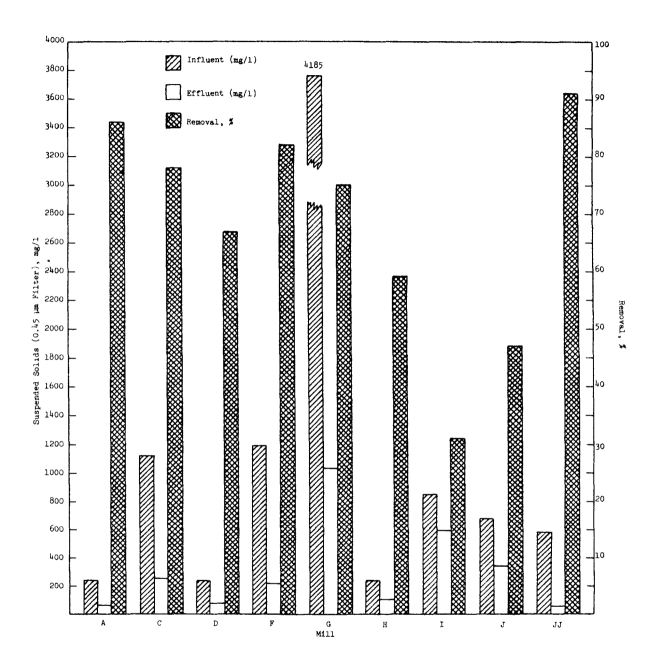


Figure 5. Suspended solids in clarifier influent and effluent collected on a 0.45- μm filter.

TABLE 8. THE VARIABILITY OF pH, BOD₅ AND COD IN CLARIFIER INFLUENT AND EFFLUENT SAMPLES FROM DIFFERENT SOURCES^{a, b}

Influent			Effluent			
рН	BOD ₅	COD	pН	BOD ₅	COD	
4.81	274	2268	6.88	268	552	
8.28	884	3899	10.68	555	1548	
10.42	482	1702	11.05	489	812	
11.53	634	2498	9.88	777 ^c	2202	
10.18	836	3388	10.80	1104°	2478	
7.35	476	1413	9.80	822 ^c	2293	
7.11	168	411	10.78	258 ^c	869	

aThese values are from all of the samples (54 sets) where pH differences and/or negative "removal" was noted.

b Influent and effluent samples were taken during the same time period.

^CProlonged holding time probably explained the apparent negative removal.

SECTION V

LABORATORY STUDY OF PROCESSES FOR HIGHER LEVELS OF SOLUBLE BOD, REMOVAL

We approached the problems of improving pulp and paper mill liquid waste treatment by two differing routes. A first route studied possibilities for improving the methods for conventional, out-plant primary clarification treatment of the total mill discharge. The second, a more innovative route, was directed to the principle of separation of flows, particularly spills of strong process water, and was also directed to individual, high-efficiency treatments, preferably within the plant, for those flows detrimental to clarifier performance.

Various treatment methods, such as sedimentation, flotation, filtration, membrane separation (reverse osmosis, ultrafiltration), carbon adsorption, and ion exchange could be used. The results of some of these methods are described in this section.

New and relatively more expensive methods of processing would be feasible for only parts of the total flow in order to prevent buildup of solubles in streams, especially those being recycled. These fractions could be treated in-plant in small, highly efficient systems or, if the streams are not to be recycled, they could be treated before they are mixed with other process waters entering the total sewer system.

However, since many of the mills involved in this study were already using clarifiers or had plans to use them, the laboratory study was carefully planned to develop a better understanding of the conventional, outplant clarification process.

SCREENING TESTS

Following the survey of mill discharges and clarifier efficiencies (Section IV), a study to optimize the removal of soluble BOD_5 from both total mill waste waters and individual streams was begun in the laboratory.

We preferred to evaluate clarification efficiency by some means other than the BOD₅ test because the 5-10 days time to run each test, depending upon the day of the week that the samples were prepared, made systematic laboratory studies very difficult. Also, we wanted a procedure based on small sample volume.

TEST TUBE TRIALS

We had developed in our laboratory a simple test tube flocculation test using a 15-ml sample in an 18×150 mm test tube mounted on a 12-inch diameter, vertically oriented, platform rotating at 3 rpm (unpublished). The

slow transfer of fluid from one end of the tube to the other produced a floc form and supernatant clarity that correlated well with jar tests with the same sample/flocculant combination. This simple, small-volume system was used in the early part of the work for screening the incoming samples.

JAR TESTS

At the same time, jar tests were carried out on the same waste material using a Phipps and Bird laboratory flocculator. Various flocculating agents, alone or in combination, were added under carefully standardized conditions of chemical addition, rapid mixing, flocculation mixing, settling and sampling. Briefly, these can be described as:

- 1. The treatment of 1.0 or 1.5 liters of waste in 3-liter jars;
- 2. The addition of the primary flocculant, followed by one minute of mixing at 100 rpm;
- 3. If a second agent (e.g., polymer, silica, activated carbon, etc.) was added, it was followed by a second rapid mix of one-half minute at 100 rpm;
- 4. The paddle speed was reduced to 20 rpm for 20 minutes to provide time for flocculation. The flocculation step was done after the final addition of chemical, no matter how many additives were involved:
- 5. At the end of the flocculation period, the jar was removed from the unit and the sample allowed to settle for one hour;
- 6. Samples were withdrawn into 250-ml bottles under slight vacuum, with the tip of the sample tube one-half inch under the surface of the liquid in the jar;
- 7. An untreated jar of the waste was carried through the entire test from rapid mix to sampling. All "removal" data are based on the comparison of the analyses of the control sample with the treated samples.

All samples were stored at 4°C (40°F) prior to analysis.

ORGANIC CARBON ANALYSIS

Use of a Beckman Process Carbonaceous Analyzer permitted relatively rapid (15 samples per day), small-volume analysis of solutions for total organic carbon (TOC). The test tube trials, after the floc was removed with the 0.45- μ m filter, provided sufficient sample for the determination of soluble organic carbon. The use of larger test tubes (25 x 250 mm) and additional sample volume on the same flocculator table, or jar tests directly, allowed determination of both total and soluble BOD₅ and TOC, and soluble organic carbon (OC) on the same sample.

We found little correlation between the reduction of soluble organic carbon and the reduction of soluble BOD $_5$ for a series of flocculation studies (Table 9). We also found no relationship of COD to BOD $_5$ during the mill survey; these results are probably due to the high ratio of nonbiodegradable to biodegradable carbon in these waste streams. Total and soluble organic carbon and BOD $_5$ components removed during flocculation would be both degradable

and nondegradable substances, with the percentage removal of degradable being overshadowed by nondegradables. While the polymers did have both biochemical and chemical oxygen demands (Table 10), the amounts added to the solutions were so small (1 to 10 mg/liter) that they did not contribute significantly to COD or BOD₅. The addition of the polymer at the 10 mg/liter level would add less than 2 percent COD and 1 percent BOD₅ to most samples if <u>all</u> of the polymer remained in solution and none was removed by flocculation and/or filtration.

TABLE 9. SCREENING TRIALS OF FLOCCULANTS AND POLYMERS BY ANALYSIS OF SOLUBLE ORGANIC CARBON AND SOLUBLE BOD5

Flocculant		Polymer	Reduct Soluble	ion ^a , % Soluble
Type	mg/1	(0.5 mg/1)	OC	BOD ₅
Alum	200	None	18.2	-9.7
Alum	200 250 250 250	Nalco 607 Nalco 627 Nalco 634 Nalco 73032	34.3 26.5 15.7 6.0	-20.9 8.1 5.4 0
FeCl ₃	100 ^b	None	20.7	20.3
	150	None	17.4	-20.5
	200	None	25.4	21.2
	200 200 200	Nalco 607 Nalco 609 Nalco 627	21.3 23.1 -7.4	16.8 20.7 15.7
	250	None	20.5	19.7
Lime	25	None	1.8	-1.9
	25	Nalco 634	-6.2	8.4
	25	Nalco 73032	2.3	0

^aValues on "as received" samples:

Soluble OC = 125-425 mg/1. Soluble BOD₅ = 100-200 mg/1.

b_{As Fe}3+

TABLE 10. COD AND BOD₅ OF VARIOUS POLYMERS USED AS FLOCCULATION AIDS

Manufacturer	Soluble COD, mg/l	Soluble BOD ₅ , mg/l
Nalco		
607	231	0
609	740	14
634	333	0
73032	402	0
Dow		
PAA	1544	214
Tydex 12	1665	0
Hercules		
Hercofloc 812.3		66

a Polymer concentration 1 g/l.

PHENOL SULFURIC ACID TEST

The phenol-sulfuric acid test (7) is a rapid method for the estimation of carbohydrates, including virtually all classes of sugars, sugar derivatives, oligo- and polysaccharides, ketoses, aldoses and aldehydes. This method has been used for several years in these laboratories for monitoring gel chromatographic columns and for the study of aldehydes in solution, both carbohydrates and noncarbohydrates (8).

Although the method appeared to be useful for measuring the removal of soluble BOD_5 during trials in the test tubes (Table 11), it did not give reproducible results in the jar tests; therefore, it was not useful as a replacement for the BOD_5 test (Table 12).

ZETA POTENTIAL

"If the zeta potential of the colloid is lowered below a critical value, the colloids tend to coalesce, resulting in coagulation" (9). Many of the "soluble" BOD $_5$ components in the waste stream are colloids, medium and low molecular weight organics, and in some cases simpler carbonaceous compounds, which should be amenable to colloidal coagulation and coprecipitation (10). Zeta potential changes and final values were compared with the removal of COD and BOD $_5$ components by flocculation (Table 13). No correlation could be drawn between the zeta potential and the reduction of either COD or BOD $_5$ for the waste stream after flocculation and filtration. There was, however, the expected correlation of near zero zeta potentials with good floc formation and supernatant clarity.

TABLE 11. COMPARISON OF PHENOL SULFURIC ACID TEST WITH BOD5
TEST TO EVALUATE FLOCCULANTS IN TEST TUBE TESTS

			Reduction,	a. _%
Floccu		Polymer	Soluble glucose	Soluble
Туре	mg/l	(1 mg/1)	equivalent	BOD ₅
FeCl ₃	50 ^b	None	28.9	22.8
	100	None	45.6	32.1
	150	None	50.0	36.2
	200	None	46.7	33.9
Lime	250	None	17.7	5.0
	300	None	6.3	18.4
	300	Nalco 609	25.6	16.4
	300	Nalco 627	40.5	12.9
	300	Nalco 634	40.5	25.9
	300	Nalco 73C32	20.0	12.9
	300	Dow PAA	39.1	23.2
	300	Dow Tydex 12	32.6	31.7
	400	None	28.1	7.1

aValues "as received":

Soluble glucose equivalent = 216 mg/l. Soluble BOD₅ = 200 mg/l.

As the study progressed we found little relation of the visual clarity of the supernatant or the rate of floc settling with the reduction of BOD_5 . A finely divided floc seemed to result in the greatest reduction in soluble BOD_5 after filtration with the 0.45- μm filter, whether or not the floc had settled.

JAR TESTS FOR OPTIMIZING BOD, REMOVAL BY CLARIFICATION

General

Following the lack of success in developing a rapid screening test to correlate flocculation with the removal of soluble BOD_5 , a number of jar tests were made with several mill effluents. We used the standard BOD_5 analysis to test these flocculants. While this resulted in a rather delayed evaluation, it could be used for: (1) a study of many flocculating agents and variables or (2) careful consideration and selection of both flocculants and process variables, with BOD_5 determinations on either (a) all samples or (b) selected samples from each series. With frequent reevaluation of the procedure as BOD_5 results became available, the (2b) scheme was selected for the majority of the jar tests.

bAs Fe³⁺.

TABLE 12. COMPARISON OF PHENOL SULFURIC ACID TEST WITH BOD 5 TEST TO EVALUATE FLOCCULANTS

IN JAR TESTS

			Redu	ction ^a , %	
Floccular Manufacturer	t mg/l	рН	Soluble glucose equivalent	Soluble BOD ₅	Soluble COD
Nalco					
607	6.0	7.5	-1.5	2.6	-1.8
609	6.0 6.0 10.0	3.0 7.5 7.5	12.0 11.5 16.5	11.3 6.8 9.8	10.2 7.2 7.2
627	6.0	3.0	4.5	8.4	5.9
634	6.0 10.0	7•5 7•5	7.3 15.3	6.8 10.0	2.7 16.6
73C32	6.0 10.0	7.5 7.5	7.3 11.5	-2.1 23.8	1.8 5.4
Dow					
PAA	6.0 10.0 20.0	7.5 7.5 5.0	15.3 15.3 10.8	2.1 12.8 -1.7	8.6 9.3 4.2
Tydex 12	6.0 6.0 6.0 10.0	3.0 5.0 7.5 7.5	3.2 -1.2 7.3 14.2	4.6 -24.8 8.1 5.1	1.3 3.3 3.8 6.8

aValues "as received":

Soluble glucose equivalent = 250 mg/l.

Soluble COD = 550 mg/l.

Soluble $BOD_5 = 220 \text{ mg/l}$.

TABLE 13. A COMPARISON OF ZETA POTENTIAL, SOLUBLE COD AND BOD5,
AND FLOC FORM IN JAR TESTS OF FLOCCULANTS

Flocc Type	ulant mg/l	Effluent ^a	pН	Zeta potential, mv	Reducti Soluble COD	on ^b , % Soluble BOD ₅	Floc ^C form
Alum	200	А		-9.3	2.1	0.0	
	395	В		0	17.3	-1.5	4+
	632	В		0	12.2	3.6	4+
Fe ³⁺	50	С		0 to -24.5	16.8	5.3	
	75	C		0	23.2	11.9	
	84	В	3.0	0	30.5	21.9	4+
	84	В	5.0	-17.8	33.3	17.2	3+
	200	Α	3.0	0	45.3	51.0	
Lime	200	A	11.6	-20.2	4.0	5.8	

^aA = special sample of beaten sulfite pulp.

A - soluble COD = 462 mg/l. soluble BOD₅ = 128 mg/l.

B - soluble COD = 453 mg/1. soluble BOD₅ = 190 mg/1.

C - soluble COD = 1284 mg/l.soluble $BOD_5 = 585 \text{ mg/l.}$

^CFloc form - 0 = supernatant clarity equal to untreated control. 5+ = clear supernatant without suspended solids.

To reduce the problem involved in setting up sampling schedules, sample transportation and storage, it was further decided to select mills by two criteria: (1) their proximity to our Appleton laboratories and (2) those having high soluble BOD₅ values in the initial survey. Others could be added, as time permitted, to allow studies of special effluents, of effluents with lower soluble BOD₅, or of effluents from more distant mills.

B = total mill effluent (Mill F).

C = digester room effluent (Mill F).

bValues for "as recieved":

TOTAL MILL EFFLUENTS WITH PRIMARY FLOCCULANTS

Tables 14, 15 and 16 summarize the large volumes of data gathered from jar tests with effluents from Mills A, D, and F, which are using conventional clarifiers or flotation systems. These data can be summarized as follows:

- 1. Ferric chloride in the range of 75-150 mg/l Fe³⁺ was the best flocculating agent for these waste streams. The optimum pH was dependent upon the particular stream being processed and the concentration of the iron salt added.
- 2. Lime, at higher concentrations (200-500 mg/l) also removed soluble BOD₅ (21-40 percent).
- 3. Alum did not remove more than 17 percent of the soluble BOD₅, although there was evidence that, with some of the waste streams, careful pH control and optimized alum concentrations could result in the removal of substantial quantities of soluble BOD₅.
- 4. The polymers plus bentonite or Celite to provide high surface area did not appear any more effective than the cheaper iron salts.
- 5. Some trials had shown that considerable amounts (22 percent) of soluble BOD₅ could be removed from the clarifier <u>effluents</u> with ferric chloride or lime (Table 14).
- 6. When the concentration of the flocculating agent and the pH are carefully controlled, from 25 to 50 percent of the soluble BOD₅ would probably be removed from the clarifier influents.
- 7. In recent similar studies in Finland ($\underline{11}$), fly ash obtained from the burning of calcium-base spent liquor was used as a flocculating agent, it was used to raise the pH during alum treatment, and it was used as a sludge conditioner. Reductions of 50 percent for BOD₅ and 70 percent for COD are reported in the treatment of kraft effluents. The authors also report that the sludge is easier to handle.

ADDITION OF ACTIVATED CARBON

Studying pilot-scale columns of activated carbon processing a chemime-chanical (CM) pulping waste, we found that 83-90 percent of the <u>total</u> BOD $_5$ was removed (12). We have no information, however, on the degree of <u>soluble</u> BOD $_5$ removed by adsorption on activated carbon. Therefore, we used jar tests to measure the reduction in soluble BOD $_5$ by powdered (300 mesh) activated carbon, with and without the addition of cationic polyelectrolytes, for the treatment of the total mill effluent (Table 17) and CM sewer discharge from Mill F (Table 18).

Soluble BOD_5 could be reduced by 32 percent in the total mill effluent and 17 percent in the CM streams. The variability of the results for the various levels of activated carbon was quite high. This was most likely due to changing conditions in the mill — high hourly variations in the flow and concentration of substances in these streams — rather than to differences in the effects of the activated carbon and polymers.

TABLE 14. JAR TESTS OF PRIMARY FLOCCULANTS WITH MILL A AND MILL D INFLUENTS AND EFFLUENTS $^{\mathbf{g}}$

			Mill A influent No additive COD = 1200 mg/l BODs = 238 mg/l	A influent dditive = 1200 mg/l = 238 mg/l	Mill A il 2 mg/l Ge COD = 1: BOD ₅ = 5	influent Gendriv 162 1178 mg/1 228 mg/1	Mill A e 2 mg/l G COD = 1 BOD 5 = 1	effluent Gendriv 162 1126 mg/l 238 mg/l	Mill D infl No additive COD = 843 BOD ₅ = 292	influent Itive 843 mg/l 292 mg/l
Floce	Flocculant Ype mg/l	Final pH	Reduction, % COD BODs	on, % BoDs	Reduction, % COD BOD	ion, % BODs	Reduction, % COD BOD	ion, % BODs	Reduction, % COD BOD;	ion, % BOD 5
None	pH study	~~~~~~ ~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	111111	11111	0000	0 0 0.0 0.0	000000 2010004	00000	8000048 600048	
F & &	70 70 70 70 70 70	6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6	31.8 31.6 23.8 23.3 7.8	14.3 10.1 11.3 11.8 8.4	23.6 26.4 27.5 27.5 26.8 26.8	21.0 25.5 13.2 19.3	23.0 23.4 26.3 24.2 6.5 23.0	22.3 21.8 10.9 12.7 20.9	22.4 21.9 22.2 21.7 21.9 23.2	11.4
Lime	100 100 100 100 100 100 100 100 100 100	8888 9.00 10.01 11.11 11.00 11	2.8 24.6 24.6	13.4 13.4 11.8 15.5	1.8 1.3 1.3 22.4 25.8		0.3 0.3 19.2 19.2	7. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1.	2.8 3.6 9.0 16.7	4.1.1.4.1.5.7.7.7.7.7.7.7.7.7.7.7.7.7.7.7.7.7.7
Solub	Soluble COD and BOD, wer	BOD 5 Wer	e measured.	14	bAs Fecls.					

TABLE 15. JAR TESTS OF PRIMARY FLOCCULANTS WITH MILL F TOTAL CLARIFIER INFLUENT⁸

Floccula		Final	$BOD_5 =$	8/13 508 mg/1 223 mg/1 ion, %		209 mg/l LO2 mg/l	$BOD_5 = 1$	9/10 440 mg/1 194 mg/1 ion, %
Type	mg/l	рН	СОД		СОД			
Lime	50 100 200 800 400 500	8.8 9.2 10.6 11.4 11.6 11.7	-7.2 0 7.2 11.8 12.9	-10.2 -10.2 -1.6 3.7 15.5	-3.8 -1.9 10.0 14.8 17.1 20.0	0 0 21.4 18.4 36.7 40.0	 	
Bentonite	50 100 150 200	 	5.1 7.6 6.5 7.6	1.4 1.0 -3.6 -1.4	 	 	 	
Fe ³⁺ (as FeCl ₃)	12.5 25.6 28.6 50.6 70.6 70.6 70.6 84.6 84.6 84.6 112.6 112.6 112.6 112.6 112.6 112.6 112.6 112.6 112.6 112.6	6.8 6.5 2.7 5.7 5.2 4.6 4.1 8.6 3.5 5.1 4.2 3.3 6.0 3.9 2.7	 		5.4 8.9 19.8 15.8 13.9 15.8	1.4 1.4 	12.9 31.3 35.8 34.8 33.7 28.0 33.3 33.7 30.5 	7.5 5.2 24.0 22.4 8.8 6.2 12.2 18.8 21.9
Alum	25 50 100 150 158b 200 250 385b 385b 385b 385b 385b 385b 385b 385b	3.5 3.5 8.9 6.8 6.2 5.3 4.6 4.3 3.9 10.6 6.9			9.0 4.3 3.8 5.2 10.0 10.0	-1.0 4.9 3.9 2.0 2.0 2.0 	8.2 8.7 19.2 21.9 21.9 23.9 19.2 11.9	3.0 -5.2 -24.2 12.0 -18.3 16.7 11.4 7.3 3.6 -2.1

aSoluble COD and BOD, were measured.
bAdded on molar basis of Fe³⁺ and Al³⁺ (0.5, 1.25, 2.0 millimolar).

TABLE 16. JAR TESTS OF POLYMERS AND ADDITIVES WITH MILL F TOTAL CLARIFIER INFLUENT

Sample 7/31 Sample 8/13 COD = 552 mg/lCOD = 508 mg/l $BOD_5 = 234 \text{ mg/l}$ $BOD_5 = 223 \text{ mg/l}$ <u>Additive</u>b Flocculant Final Reduction, % Reduction, % COD Manufacturer mg/lmg/l $_{\mathrm{Hq}}$ COD Туре BOD_5 BOD₅ Nalco 607 1.0 Bentonite 200 8.2 11.9 607 A/C 1.0 200 7.0 -4.6 -----607 6.0 -1.8 None 7.5 2.6 609 1.0 Bentonite 200 9.4 3.2 __ 609 1.0 A/C 200 8.2 18.7 6.8 7.2 609 6.0 None 7.5 609 6.0 None 3.0 10.2 11.3 10.0 7.5 7.2 9.8 609 None 627 1.8 1.0 Bentonite 200 9.2 A/C 627 1.0 200 __ ---___ 9.6 16.9 8.4 6.0 3.0 5.9 627 None __ 634 200 8.0 -1.41.0 Bentonite 634 1.0 A/C 200 1.4 __ 5.1 634 6.0 None 7.5 2.7 6.8 634 10.0 None 7.5 10.0 16.6 6.7 23.8 7.4 73C32 1.0 Bentonite 200 9.1 73032 1.0 A/C 200 __ 8.0 19.6 1.8 73032 6.0 None 7.5 -2.1 --73C32 10.0 None 7.5 5.4 23.8 6.1 29.4 73032 10.0 Celite 200 7.5 ---3.0 21.2 Dow Tydex 12 1.0 Bentonite 200 12.4 -1.4--Tydex 12 1.0 A/C 200 9.0 20.5 __ Tydex 12 6.0 None 7.5 3.8 8.1 ----Tydex 12 6.0 None 5.0 3.3 -24.8 Tydex 12 6.0 None 3.0 1.3 4.6 6.8 Tydex 12 5.1 10.0 None 7.5 PAA 1.0 Bentonite 200 14.7 -4.5 8.6 A/C __ __ -11.4PAA 1.0 200 8.6 PAA 6.0 None 7.5 2.1 ----6.3 21.8 PAA 10.0 None 7.5 9.3 12.8 PAA 10.0 Celite 200 7.4 14.2 4.2 PAA -1.720.0 None 5.0 ___

^aSoluble COD and BOD₅ were measured.

bA/C = activated carbon (Filtrasorb 300, powdered).

TABLE 17. JAR TESTS OF THE EFFECT OF ACTIVATED CARBON AND VARIOUS ADDITIVES ON SOLUBLE COD AND BOD5 IN MILL F CLARIFIER INFLUENT

(1) 50 None	Acti	vated					8/13 508 mg/l 223 mg/l		8/21 209 mg/1 102 mg/1		9/10 26 mg/1 196 mg/1
Type ^a mg/l Type mg/l pH COD BODs COD BODs COD E (1) 50 None	car	rbon	Additiv	96	Final	Reduct	ion. %	Reduct	ion. %	Reduct	ion. %
100 None	Type ^a	mg/l									BOD ₅
100 None	(1)	50	None		7.4	6.1	19.2	4.1	3.4		
200 None		100	None		7.5	5.7	8.7	6.9			
200 Nalco 73C32 1.0 7.5 8.0 19.6 6.3 2 200 Dow Tydex 1.0 7.4 9.0 20.5 5.2 200 Cellulose 500 5.2 1 200 Nalco 609 1.0 7.9 8.2 18.7 200 Nalco 627 1.0 7.9 9.6 16.9 200 Nalco 634 1.0 7.8 5.1 1.4 200 Dow PAA 1.0 7.9 8.6 (2) 50 None 14.7 17.0 100 None 17.4 7.5 200 Nalco 73C32 1.0 11.9 27.4 200 Dow Tydex 1.0 16.4 1 200 Cellulose 500 8.0 1 200 Cellulose 500 8.0 1 200 Cellulose 500 5.2 1		150	None		7.5	5.9	12.3				
200 Dow Tydex 1.0 7.4 9.0 20.5 5.2 200 Cellulose 500 5.2 1 200 Nalco 609 1.0 7.9 8.2 18.7 200 Nalco 627 1.0 7.9 9.6 16.9 200 Nalco 634 1.0 7.8 5.1 1.4 200 Dow PAA 1.0 7.9 8.6 200 None 17.4 7.5 200 None 17.4 7.5 200 None 11.9 27.4 200 Nalco 73C32 1.0 11.9 27.4 16.4 1 200 Dow Tydex 1.0 16.4 1 200 Cellulose 500 8.0 1 200 Cellulose 500 4.0 1 4.0 1 200 Cellulose 500 5.2 1		200	None		7.4	5.7	-5.5	13.4	11.5	3.5	2.0
200 Cellulose 500 5.2 1 200 Nalco 609 1.0 7.9 8.2 18.7 200 Nalco 627 1.0 7.9 9.6 16.9 200 Nalco 634 1.0 7.8 5.1 1.4 200 Dow PAA 1.0 7.9 8.6 (2) 50 None 14.7 17.0 100 None 17.4 7.5 200 Nalco 73C32 1.0 11.9 27.4 200 Dow Tydex 1.0 16.4 1 200 Cellulose 500 8.0 1 200 Cellulose 500 4.0 1 + Tydex 1.0 5.2 1		200	Nalco 73C32	1.0	7.5	8.0	19.6			6.3	23.5
200 Nalco 609 1.0 7.9 8.2 18.7 200 Nalco 627 1.0 7.9 9.6 16.9 200 Nalco 634 1.0 7.8 5.1 1.4 200 Dow PAA 1.0 7.9 8.6 200 None 17.4 7.5 200 None 11.9 27.4 16.4 1200 Dow Tydex 1.0 16.4 1200 Cellulose 500 8.0 1200 Cellulose 500 8.0 1200 Cellulose 500 4.0 1200 Cellulose 500 5.2 1200 Cellulose 500 5.2 1		200	Dow Tydex	1.0	7.4	9.0	20.5			5.2	6.1
200 Nalco 627 1.0 7.9 9.6 16.9 200 Nalco 63\(^4\) 1.0 7.8 5.1 1.\(^4\) 200 Dow PAA 1.0 7.9 8.6		200	Cellulose	500						5.2	12.2
200 Nalco 634 1.0 7.8 5.1 1.4		200	Nalco 609	1.0	7.9	8.2	18.7				
200 Dow PAA 1.0 7.9 8.6 (2) 50 None 14.7 17.0 100 None 17.4 7.5 200 None 11.9 27.4 16.4 1 200 Dow Tydex 1.0 16.4 1 200 Cellulose 500 8.0 1 200 Cellulose 500 4.0 1 + Tydex 1.0 4.0 1 200 Cellulose 500 5.2 1		200	Nalco 627	1.0	7.9	9.6	16.9				
(2) 50 None 14.7 17.0 100 None 17.4 7.5 17.0 None 11.9 27.4 16.4 1 200 Nong 1.0 16.4 1 200 Cellulose 500 8.0 1 200 Cellulose 500 4.0 1 + Tydex 1.0 4.0 1 200 Cellulose 500 5.2 1		200	Nalco 634	1.0	7.8	5.1	1.4				
100 None		200	Dow PAA	1.0	7.9	8.6					
200 None	(2)	50	None		<u></u>			14.7	17.0		
'200 Nalco 73C32 1.0 16.4 1 200 Dow Tydex 1.0 7.5 200 Cellulose 500 8.0 1 200 Cellulose 500 4.0 1 + Tydex 1.0 5.2 1		100	None					17.4	7.5		
200 Dow Tydex 1.0 7.5 200 Cellulose 500 8.0 1 200 Cellulose 500 4.0 1 + Tydex 1.0 5.2 1		200	None					11.9	27.4		
200 Cellulose 500 8.0 1 200 Cellulose 500 4.0 1 + Tydex 1.0 200 Cellulose 500 5.2 1		1200	Nalco 73C32	1.0						16.4	12.2
200 Cellulose 500 4.0 1 + Tydex 1.0 4.0 1 200 Cellulose 500 5.2 1		200	Dow Tydex	1.0						7.5	9.2
+ Tydex 1.0 200 Cellulose 500 5.2 1			Cellulose	500							11.2
200 Cellulose 500 5.2 1		2 0 0	Cellulose	500						4.0	19.4
			+ Tydex	1.0							
+ 73032 1 0		200	Cellulose	500						5.2	14.3
ve e			+ 73032	1.0							
300 None 17.9 13.2		7	None					17.9	13.2		
400 None 17.9 32.1		400	None					17.9	32.1		

a(1) Filtrasorb 300, powdered.

In all of the jar tests the samples were first screened through a 60-mesh stainless steel screen, the activated carbon (and polymers, if used) was added, and the sample was processed through the mixing and settling cycles.

No attempt was made to correlate the degree of soluble BOD₅ removal with the presence of fiber fines or other carbonaceous adsorbing materials.

Although the activated carbon-soluble BOD₅ complex could be readily filtered by a 0.45-µm filter, it did not settle in the jars with or without the use of polymer flocculants. Activated carbon, therefore, seems useful as an additive to the primary system for the reduction of BOD₅ if the carbon could be removed in either the primary or secondary system. Larger carbon particles might be easier to remove; however, they would have less surface area and, therefore, probably be less sorptive.

EFFECT OF THE CONCENTRATION OF THE EFFLUENT

If the total water use in the mill were reduced through recycling, the wastes would be more concentrated. In order to study the flocculation and soluble BOD₅ removal characteristics of such wastes we concentrated by reverse osmosis a sample of the total mill effluent (clarifier influent) from

⁽²⁾ Same alcohol and water washed (oven dried).

the Mosinee mill. We have been evaluating and developing uses for reverse osmosis (RO) in our laboratories since 1964 ($\underline{13}$). This process can concentrate solutions and at the same time remove some of the soluble BOD₅.

TABLE 18. JAR TESTS OF THE EFFECT OF ACTIVATED CARBON AND ADDITIVES ON COD AND BOD₅ OF MILL F CHEMIMECHANICAL PULPING EFFLUENT^{8.}

Carbon, b	Additive		Reduct		Final
mg/l	Type	mg/l	COD	BOD ₅	ДЩ
50	None		2.5	15.9	7.4
100	None		3.4		7.5
125	None	word visual	5•9	11.6	7.6
150	None		4.5	5.2	7.7
200	None		5.2	12.0	7.5
200	Nalco 634	10.0	4.3	2.0	7.5
200	Nalco 73032	10.0	5.9	13.1	7.5
200	Dow Tydex 12	10.0	10.0	9.0	7.5
200	Dow PAA	10.0	10.5	16.8	7.5
300	None		10.2	9.8	7.5
None	Nalco 634	10.0	-1.0	-1.5	7.4
None	Nalco 73C32	10.0	1.0	5.9	7.4
None	Dow PAA	10.0	-4.1	-5.1	7.6

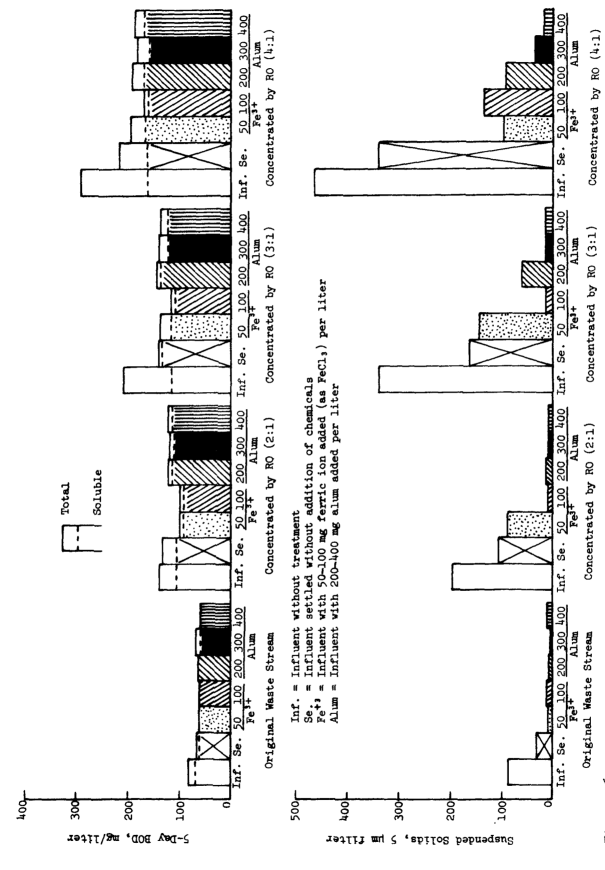
a Soluble COD and BOD, were measured.

Initial values: Soluble COD = 1105 mg/l Soluble BOD₅ = $\frac{190 \text{ mg}}{1}$.

Reverse osmosis was used to make a fourfold v/v concentrate of the total mill effluent. This solution was diluted with the permeate (fluid passing through the RO membrane during the concentration process) to prepare 3:1, 2:1, and 1:1 concentrations. Jar tests were made with all concentrations, and the samples were analyzed for suspended solids, total and soluble BOD₅, and color before and after treatment with ferric chloride-polymer and alum-polymer combinations (Figure 6).

These data indicate that treatment with ferric chloride at 100 mg Fe³⁺ per liter and Hercofloc 812.3 at 0.75 mg per liter could produce an acceptable effluent for discharge, in terms of suspended solids and BOD₅, up to concentration of approximately 3:1 and reduce both hydraulic and gravimetric loads.

Activated carbon - Filtrasorb 300, powdered, alcohol/water washed, oven dried.



Suspended solids and BODs in various concentrations of influents treated with Fe3+ or alum. Figure 6.

With effluents more concentrated than 3:1, flocculating agents did not produce adequate effluents.

While this was a rough study and practical means of processing were not worked out — we did not study spill control, the effect of spills on water reuse within the mill, or economics — it certainly indicates that wastes can be successfully treated at higher concentrations.

JAR TESTS ON WASTE STREAMS PRIOR TO PILOT SCALE TRIALS

Mosinee Paper Company of Mosinee, Wisconsin and the Combined Locks mill of Appleton Papers Division of NCR, Combined Locks, Wisconsin were selected for the two pilot-scale clarifier trials. Jar tests were made on the total mill effluents. These preliminary tests were followed periodically by other jar tests while the field trials were in progress.

Tests on Mosinee Paper Company Effluent

Samples from these jar tests were, in some cases, analyzed completely for suspended solids, total and soluble BOD₅, organic carbon, COD and color (Tables 19 and 20) in order to measure the removals that might be attained under different conditions of treatment. Others were merely used to establish flocculating characteristics (Table 21) and/or sludge volumes that would develop with the addition of the chemical combinations.

In the early trials we established that alum at 200-300 mg/l and ferric chloride at 100 mg/l Fe³⁺ could be used in jar tests to remove BOD₅ - 35-55 percent (total) and 12-25 percent (soluble) - from the wastes. The addition of 0.75 mg/l Hercofloc 812.3 appeared to provide additional removal of soluble BOD₅: removal increased from 13 to 35 percent when Hercofloc was added with 100 mg/l alum, and from 12 to 28 percent with 200 mg/l alum (Table 19). The use of the Hercofloc polymer alone at 0.75 and 1.5 mg/l markedly increased the removal of soluble BOD₅: we found 14 and 21 percent removal, respectively.

Lime as a primary flocculating agent and activated silica as an additive (Table 20) appeared to be relatively effective (4-16 percent) in the removal of soluble BOD₅ from the Mosinee mill effluent.

Of great interest in these studies was the high removal of COD, color, and total and soluble BOD₅ that could be achieved with ferric chloride and alum from the clarifier <u>effluent</u> from this mill (Table 22). Twenty-five to forty percent removal of the soluble BOD₅ components was obtained with these primary flocculants added to either the north or south clarifier effluents. Lime and "additional" Hercofloc 812.3 did not enhance the removal of the "solubles" but did increase the removal of total BOD₅, COD and color.

Adjustment of the pH from the initial ll, or higher, to 8 with sulfuric acid prior to the addition of the chemicals, markedly increased the amount of suspended solids removed. This only slightly affected the amount of soluble BOD₅ remaining in solution (Table 23). The ferric ion was equally effective as the chloride or sulfate salt when they were used at the same concentration

TABLE 19. JAR TESTS OF THE EFFECT OF FLOCCULANTS ON BOD5, COD, ORGANIC CARBON, AND COLOR IN MOSINEE CLARIFIER INFLUENT

				%]	Reducti		
Floccu	lant	Final	Tot	al	Sol	_uble_b	Total
Type	mg/l	рН	BOD ₅	COD	OC	BOD ₅	color
FeCl ₃	50 ^d 100 150	6.4 6.0 5.0	51 37 58	71 70 76	31 44 47	13 25 24	52 43 89
Alum	100 200 300	7.2 7.0 6.8	49 56 56	60 68 73	14 24 31	13 12 12	24 43 47
Lime	200 300 500	11.1 11.6 12.0	22 39 41	52 68 71	-13 -1 1	-16 -14 -9	15 58 11
Hercofloc 812.3	0.75 1.5	8.4 8.2	18 19	8 11	3 6	14 21	0 15
Hercofloc 8 following p			as added	after	l min	mixing	with the
FeCl ₃	10 25 50	7.0 6.8 6.4		oc formoc form		30	47
Alum	50	7.4	No fl	oc for	nation		

54

52

76

71

14

21

35

28

18

32

7.2

7.0

Total COD = 711 mg/l

100

200

 $BOD_5 = 215 \text{ mg/l}$

Color = 109 units. Soluble OC = 87 mg/l

 $BOD_5 = 122 \text{ mg/l}$

a Initial control values for sample taken 11/21/74:

 $[^]b\mathrm{Filtered}$ through a 0.45 µm Metricel filter. $^c\mathrm{Co-Pt}$ units at 465 nm (NCASI Method). $^d\mathrm{As}~\mathrm{Fe}^{3+}.$

TABLE 20. JAR TESTS OF FeCl3, ALUM, AND ADDITIVES AS FLOCCULANTS FOR MOSINEE CLARIFIER INFLUENT

Floor	ulant				Redu	ction	, %	
FIOCU	uranc	b /2	Final	Tot	ta.l	So.	luble_	
Primary, mg/l	Polymer	, mg/l	pН	BOD ₅	COD	oc	BOD ₅	Color
Analytical grachemicals:	de prima	сy						
FeCl₃ 50 ^c	Herc.	0.75 1.5	5.8 5.8 5.8	38 37 38	56 57 58	49 51 51	29 27 27	88 89 90
Alum 50	Herc.	0.75 1.5 10	7.4 7.5 7.7 7.7	11 10 6 5	-0.7 -0.7 0.7 4	20 21 14 24	9 15 10 4	39 47 45 40
100	Herc.	0.75 1.5 10	7.1 6.8 7.1 7.4	13 15 10 11	3 16 3 4	34 33 34 29	5 8 16 12	69 69 72 53
200	Herc.	0.75 1.5	7.4	23 23 21 23	54 54 47 57	42 40 39 39	8 4 4 4	86 88 87 84
Commercial grachemicals:	de prima	ry				-		
FeCl ₃ 50 ^c	Herc.	0.75	5•5	32	50		28	
Alum 100	Herc.	0.75	6.5	20	9		13	

a Initial values (for untreated influent control) for sample taken 12/12/74:

Total COD = 711 mg/1

 $BOD_5 = 215 \text{ mg/l}$

Color = 109 Pt-Co units

Soluble OC = 87 mg/lBOD₅ = 122 mg/l.

Herc., Hercofloc 812.3; Act.Sci., activated silica.

CAs Fe³⁺.

of Fe³⁺ and when the pH of the influent was not above 9.0 (Figure 7). Sludge volumes, determined with separate jar tests and the Imhoff cone, were also markedly reduced by the addition of the Hercofloc 812.3 (Table 24) and for this reason it was included in <u>all</u> but one of the studies at this mill.

TABLE 21. JAR TESTS OF THE EFFECTS **OF** PRIMARY FLOC**C**ULANTS
AND POLYMERS ON pH AND CLARITY OF
MOSINEE CLARIFIER INFLUENTS

	Floc	culant				Influe	nt samples		
Prima	rv.	Polym	er.	1,	/16/75	3	/5/75	3,	/11/75
mg/		mg/		Нq	Claritya	Hq	Claritya	pН	Claritya
Fe ³⁺	25							6.8	0
10	50			6.6	0	6.9	0	5.9	5
) •	Herc.	0.75	6.8	Ö	6.6	Ö	/·/	
	100			5.0	Ö	5.7	4	4.2	4
	100	Herc.	0.75	5.6	3	5.6	λ,	4.3	14
	150				- -	4.5	i	3.4	3
		Herc.	0.75			4.3	ī	3.4	3 3
Alumb	100					9.0	0	~	
		Herc.	0.75			8.6	0		
	200					8.4	1	****	
	300					7.2	14		
Alum ^c	200			7.4	1				
111 0011	300			7.1	5		= -		
	400			6.5	Ĺ				
d									
Alum	200			7.0	1				
	300			6.9	5				
	400			6.5	4				

aClarity: 0 = untreated control

Tests on Combined Locks Mill Effluent

Since Mill F had been one of those in the preliminary jar test studies on the optimization of soluble BODs removal, only a few additional tests were conducted prior to the pilot-scale field trials. Data from these tests (Table 25) substantiated the earlier findings (Tables 15 and 16) that ferric chloride, ferric sulfate and alum were all effective in removing total and soluble BODs from this waste stream. We found, however, marked variations in the amounts of removal with the different concentrations and combinations of the flocculants used, apparently depending upon the amounts of spent cooking liquor in the mill effluent.

^{1-5 =} partial to complete clearing of the supernatant after
 one hour settling.

Analytical grade.

Commercial grade with precipitate in suspension.

Commercial grade decanted to remove the precipitate.

TABLE 22. JAR TESTS OF THE EFFECT OF FLOCCULANTS ON pH, BOD₅, COD, AND COLOR OF EFFLUENTS FROM MOSINEE CLARIFIERS^a

					luction, %	
Floccul				Total		Soluble
Туре	mg/l	pН	BOD ₅	COD	Color	BOD ₅
From Nort	th Clarifi	ler:				
FeCl ₃	50 ^b 100 150	5.1 3.4 3.0	61 63 64	72 72 68	65 79 85	39 40 38
Alum	100 200 300	6.0 6.2 6.2	43 61 61	56 70 70	42 65 68	26 25 26
Lime	200 300 500	9.9 11.1 11.6	25 33 41	32 48 55	28 42 42	-14 -19 0.8
Hercofloo 812.3	0.75 1.5	8.8 8.6	12 15	4 11	7 4	- 6 3
From Sou	th Clarif	ier:				
FeCl ₃	50 ^b 100 150	5.7 3.8 3.1	63 65 59	68 70 64	46 70 75	38 39 38
Alum	100 200 300	6.9 6.5 5.6	No f 60 62	locculat 64 66	ion 27 46	31 26
Lime	200 300 500	11.3 11.6 12.0	No f 43 43	locculat 44 50	ion 2 11	- 13
Hercofloo 812.3	0.75 1.5		No f	locculati	Lon	

 $^{^{\}rm a}$ Initial values (control) of samples taken 11/21/74:

Soluble BOD (mg/l) North=124 South=130

Total BOD (mg/l) North=214 South=240

COD (mg/l) North=628 South=612

Color (units) North=628 South=612 bAs Fe³⁺.

TABLE 23. EFFECT OF pH ON JAR TEST FLOCCULATION OF MOSINEE CLARIFIER INFLUENT

	flocculant	Final pH	BOD ₅ red Total	uction, % Soluble	Reduction in suspended solids,
Fe ³⁺	50 100 150 200	10.7 10.3 9.9 9.4	-3 -5 1	-5 -3 6 8	22 55 72 50
Alum	200	10.8	-0.2	1	35
	300	10.6	-7	4	18
	400	10.4	4	12	2
	500	10.2	2	7	15
	600	10.1	2	5	30
Influent	t adjusted 1	to pH 8 wi	th sulfurio	acid prior	to jar test:
Fe ³⁺	50	7.2	6	11	75
	100	6.8	6	4	72
Alum	200	7.5	-9	5	76
	300	7.3	-5	- 7	84
	400	7.1	-8	20	80

 $^{^{}m a}$ Influent sample of 2/11/75 had an initial pH of 11.2.

The effect of these spent liquors and other mill discharges on the overall efficiency of the clarifier is discussed in Section VII.

LABORATORY SURVEY OF SOURCES OF SOLUBLE BOD, IN MILL SYSTEMS

In a mill system with multiple sewers there are certain streams that carry the bulk of the BOD and COD. Some of the BOD and COD can be more easily removed from concentrated streams than from diluted waste, and some substances may alter the properties of others or interfere with their removal. Therefore, isolation and treatment of these more concentrated streams would be a logical way to improve the removal of BOD₅ by the clarifiers and thereby reduce the BOD₅ load on the out-plant treatment system. Importantly also, the degree of recycling of the process water could be substantially increased.

Separate Sewer Discharges

In our study samples from the separate sewer discharges were obtained from two of the mills and analyzed for various types of solids, for soluble and total BOD₅, as well as soluble and total COD components (Tables 26 and 27).

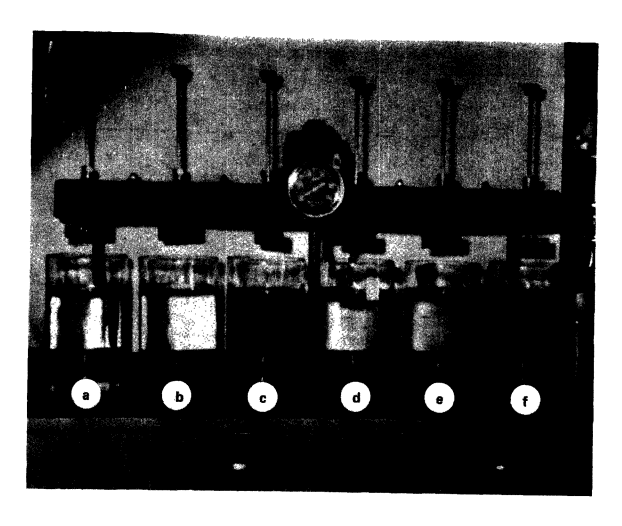


Figure 7. In jar tests with Mosinee mill effluent, pH 8.5, ferric chloride and ferric sulfate at 25 and 50 mg/l of Fe³⁺ (a), (b), (d), and (e), precipitated more solids than when they were used at a higher concentration, 100 mg/l of Fe³⁺ (c) and (f).

If one or more of the high BOD₅ streams (the discharges from the CM sewer in Mill F or the pulp mill, soda recovery or fifth stage brownstock washer sewers in Mill K) could be treated separately by flocculation, reverse osmosis, ultrafiltration or some other process, the overall efficiency of the treatment plant would increase, there would be less of a load on secondary facilities, and there would be a greater potential for water reuse (details discussed in Section VII).

Breakdown of Fibers

Another source of soluble BOD_5 was fibers: saccharides and other microbial nutrients are released during the beating of pulp or during the recycling of fibers in water recovery systems.

In order to study this release of soluble BOD₅, a beater test was arranged with a sample of unbleached sulfite pulp in a laboratory beater using

a conventional 1-1/2 pound (0.68 kg) test with 12 pounds (5.5 kg) on the bed-plate. One-liter samples were removed at half-hour intervals during the 90-minute test period and freeness, total solids, total and soluble BOD_5 , and total and soluble COD were analyzed on each sample (Table 28).

TABLE 24. VOLUME OF SLUDGE FROM MOSINEE CLARIFIER INFLUENTS TREATED WITH VARIOUS FLOCCULANTS AND POLYMER

	Floc	culant		Sludge volume,	Final	Clarityb
Primary,	mg/l	Polymer	a mg/l	ml/l	рН	0-5
Influent	12/21/	74:				
Fe ³⁺	50 50	None Herc.	 0.75	122 115	5.4 5.5	4 5
Alum	100 100	None Herc.	<u></u> 0.75	64 54	6.2 6.5	2 2
Influent	12/31/	74:				
Fe ^{3+C}	50 50	None Herc.	<u></u> 0.75	92 80	6.3 6.2	4 5
Fe ^{3+d}	50 50	None Herc.	 0.75	96 85	6.3 6.3	5 5
Influent	1/16/7	5:				
Fe ³⁺	50 50 100 100	None Herc. None Herc.	0.75 0.75	36 30 48 60	6.6 6.8 5.0 5.6	0 0 0 3
Alum	300 300	None Herc.	- - 0.75	96 80	6.3 6.3	4 5

Herc., Hercofloc 812.3.

Soluble BOD₅ and soluble COD markedly increased as the beating progressed and the fibers were fibrillated, releasing materials which became microbial nutrients in the BOD₅ test.

We ran several flocculation tests on portions of the filtrate from a 90-minute beaten sample (D in Table 28). This sample was passed through a 0.45- μ m filter to remove fibers and insoluble BOD₅ and COD components. Ferric chloride was again the best "single" flocculating agent (51 percent removal

bClarity: 0 = control (no apparent flocculation)

^{5 =} clear supernatant.

CAnalytical reagent grade of FeCl₃. dCommercial grade, 42°Be, of FeCl₃.

TABLE 25. REMOVAL OF BOD, FROM EFFLUENTS OF COMBINED LOCKS MILL BY FLOCCULANTS AND POLYMERS

	Floccula	ınt	· · · · · · · · · · · · · · · · · · ·	Efflue BOD ₅ re	ent 4/23 ^b	Efflue BOD ₅ re	ent 4/24 ^c
Primary,	mg/l	Polymer,	a mg/l	Total	Soluble	Total	Soluble
FeCl ₃ d	10 25	None None			cculation cculation		
	50 50 50 50 75	None H812.3 N73C32 None	0.75 1.0	 38.3	 30.9	18.9 18.5 30.6 29.7	15.5 14.4 26.3 20.1
	75 75 100	N73C32 H813.2 None	1.0 0.75	36.9 42.0	21.8	27.5 26.1	23.7 18.6
Fe ₂ (SO ₄) ₃ ^d	25 37·5 50 50 50 75 75	None None None H812.3 N73C32 None H812.3	 0.75 1.0 0.75 1.0	No flo	eculation eculation cculation 20.6 	36.9 26.6 29.3 22.5 38.7 32.0	27.8 16.0 20.1 9.8 28.4 35.6
Ferri-floc ^d	75 75	н812.3 N73C32	0.75 1.0	,		26.6 20.7	14.4 15.5
Alum	50 100 200 300	None None None	 		eculation eculation 	8.1 25.2	13.9 10.3
Lime	200 200	н812.3 N73C32			cculation cculation		
None		н812.3	0.75			5.4	-6.7

aH812.3, Hercofloc 812.3; N73C32, Nalco 73C32. b4/23 BOD₅: Total = 274 mg/l

Soluble = 220 mg/l.

c_{4/24} BOD₅: Total = 222 mg/l Soluble = 194 mg/l.

d_{As Fe³⁺.}

TABLE 26. SOURCES OF SOLUBLE AND TOTAL COD AND BOD, IN EFFLUENTS OF MILL Fa

					Solids				COD			BODs	ı
Source	Flow, mgd	Ηď	Suspended mg/l %Tot	ended % Total	Diss mg/1	Dissolved mg/l % Total	Total, mg/l	Rg/1	Soluble mg/1 % Total	Total, mg/l	Sol mg/l	Soluble mg/l % Total	Total, mg/l
Main sewer	. ო	7.63	860	51.9	962	1.84	1656	309	4.62	1052	119	4.74	251
CM sewer	1.3	7.75	2322	59.5	1598	40.8	3920	1284	8.69	1838	585	100.5	582
Wrapper room	1.2	7.58	620	61.1	394	38.9	1014	172	17.1	1006	35	24.0	146
No. 6 sewer	0.95	7.60	1618	6.69	869	30.1	2316	122	8.8	1394	28	15.5	181
Clarifier influent	ቱ.7	7.63	972	16.0	1140	54.0	2112	453	30.5	1486	190	52.9	359
0													

aSamples taken 9/10/74.

TABLE 27. CHEMICAL, BIOLOGICAL, AND PHYSICAL ANALYSES OF VARIOUS PROCESS STREAMS WITHIN THE MOSINEE MILL (MILL K)

Source	ЪН	Total, mg/l	mg/1 BOD ₅	Soluble	Soluble, mg/la OC BODs	Colorb	Suspended solids 0.45 µm 5 µm	solids 5
Digester sewer	11.8	2971	635	747	775	0001	239	98.5
Pulp mill sewer	11.2	0944	1624	1099	1074	4880	260	71
5th Stage brownstock washer	10.7	4865	1349	1402	1156	4150	130	75
High density sewer	5.6	553	158	138	1441	920	120	56
Foul evaporator	8.5	871	762	197	624	100	22	34
Digester blowdown	. c.	1485	1 796	389	953	562	ત	1.5
Soda recovery sewer	10.1	8328	4162	2077	3705	4200	989	184
Clarifier influent	8.0	920	181	47	106	92	1004	989

aPassing through a 0.45 µm Metricel filter. bCo-Pt units at 465 nm (NCASI Method). Collected on filters of 0.45 and 5 µm pore size.

of soluble BOD₅) (Table 29). We added cellulose powder (500 mg/l, Whatman, Standard grade) to provide particles of high surface area and enhance floculation (No. 3 series in Table 29); it slightly increased the removal of soluble BOD₅ (1-9 percent). At pH 7.7 with alum, the removal was 2 percent better than at pH 4.2 (2A versus 1A). The addition of cellulose powder and a pH of 7.7 further increased (to 9 percent) the soluble BOD₅ that was removable with 200 mg/l alum. A combination of cellulose, lime and alum at pH 10.7 (4C) was 23 percent better than lime alone at pH 11.8 (1C) or lime and alum without cellulose (2C) at pH 10.7 (29 percent increase). Additional work on the effect of these high BOD₅ streams on the efficiency of clarifiers was done later in the study and is discussed in greater detail in Section VII.

TABLE 28. RELEASE OF SOLIDS, SOLUBLE COD, AND BOD, BY BEATING.

			24 Hr	Solu	uble COD	Solu	uble BOD ₅
Sample no.	Beating, min	Freeness,	solids, g/l	mg/l	Increase,	mg/l	Increase,
A	0	750	0.29	334		77	
В	30	370	0.32	396	18.6	77	0
C	60	100	0.35	416	24.6	118	53.2
D	90	30	0.38	462	38.3	128	66.2

al-1/2 1b beater method with 5.5 kg load on the bed plate.

ALTERNATE METHODS FOR REDUCTION OF BOD5

Ultrafiltration is a process for the separation of soluble low and high molecular weight (or size) components into two separate streams through the use of semiselective membranes under low pressure (less than 300 psig or 21 kg/sq cm). We used ultrafiltration in two trials with effluents from Mill F.

In the first trial a sample of the total mill discharge to the clarifier was processed through a Westinghouse module using a polysulfone membrane that had high pH and temperature limits (pH 11.5 and 95°C). Removal of 59-65 percent of the soluble BOD_5 was attained in the permeate (that portion of the solution passing through the membrane). The permeate apparently contained low molecular weight components, such as acetic acid, that had not been removed by flocculating agents (Table 30, Part A). The flocculating agents, lime with the polymers Nalco 609 or Dow PAA (polyacrylamide) did not remove additional soluble BOD_5 from the permeate of the total mill effluent.

The second trial (Table 30, Part B) was conducted with a sample of the CM sewer effluent, which had been shown in previous work to contain 33 percent of the soluble BODs in 18 percent of the flow going to the clarifier.

^DCanadian standard.

Ultrafiltration without additives removed 69 percent of the soluble BOD_5 , and activated carbon did not remove any more BOD_5 . Apparently the membrane was the controlling factor in the removal of soluble BOD. The carbon did not adsorb any of the transferrable components probably because its adsorptive capacity was used up by larger, more heavily charged molecules.

TABLE 29. JAR TESTS OF VARIOUS FLOCCULATION TREATMENTS
OF FILTERED 90-MINUTE BEATER SAMPLES^a

					So	luble COD	Sol	ıble BOD ₅
Tre	eatment and	Floccu				Reduction,		Reduction,
	sample	Туре	mg/l	pН	mg/l	% 	mg/l	%
1.	Flocculant added:							
	Α	Alum	200	4.2	458	2.1	105	0.0
	В	FeCl ₃	200	2.7	256	45.3	51	51.0
	C	Lime	200	11.8	449	4.0	98	5.8
2.	Above sampl treated wi alum and flocculate	th						
	A	Alum	200	7.7	336	28.2	102	1.9
	C	Alum	200	10.7	449	4.0	109	0.0
3.	Cellulose r (500 mg/l) to No. 1 s	added						
	A	Alum	200	4.2	400	14.5	95	8.6
	В	FeCl ₃	200	2.7	248	47.0	50	51.9
	C	Lime	200	11.8	422	9.8	90	13.5
	D	Control	None		468		104	
4	Alum added t No. 3 sampl							
	A	Alum	200	7.7	355	22.0	88	15.4
	C	Alum	200	10.7	406	13.2	74	28.8

a Initial values: pH = 6.7

Soluble COD = 462 mg/l

Soluble BOD₅ = 128 mg/l.

TABLE 30. SOLUBLE COD AND BOD₅ IN MILL F EFFLUENTS TREATED WITH FLOCCULANTS AND REVERSE OSMOSIS OR ULTRAFILTRATION

	ī	Flocculants	ants	Membrane	,	Sol	Soluble COD Reduction,	Soluk	Soluble BODs Reduction,
	Stream	Type	mg/1	treatment	Sample	mg/1	9	ıng/⊥	%
Α.	Clarifier influent	None	1	UF	Feed Permeate	433 136	68.6	220 78	64.5
		Nalco 609	300	UF	Feed Permeate	382 138	63.9	180	 1.65
		Dow PAA	10.0	UF	Feed Permeate	433 138	68.1	206 73	9.49
B	CM sever	None Activated	1	UF	Feed Permeate	1095 292	73.3	528 162	69.3
		carbon	200	UF	Feed Permeate	901	75.8	426 137	67.8
		None	1	UF + RO	Feed Perm. UF Perm. RO	1095 392 58	 64.2 94.7	506	92.5
		None	1	RO	Feed Permeate	1196	91.8	522 57	 89.1
		None None A/C	500	UF + RO	Feed Perm. UF Perm. RO	1039 306 36	70.5	489	93.7
		FeCl ₃	100 (as Fe ³⁺)	UF	Feed Permeate	795 297	62.6	508 180	9.49

The ultimate value of ultrafiltration might be in concentrating the high molecular weight soluble BOD₅ components and removing them from process streams. It has been demonstrated elsewhere (21) that ultrafiltration could be carried to a concentration of 25-35 percent solids without complications from osmotic pressure buildup which directly affects RO flux rates and concentrations. This would be sufficiently high for final disposal by discharge into the strong pulping liquor streams for evaporation and utilization or burning. Still another use for ultrafiltration is in reducing the buildup of high molecular weight solubles in recycled process water within the mill; this could reduce water consumption and the volume of overflow to the outplant clarifier.

Reverse osmosis, similar to ultrafiltration but operating at higher pressures (400-800 psig or 28-56 kg/sq cm) with tighter membranes which reject smaller molecules, was tried both as a possible treatment method and as a further refinement for the permeate from the ultrafiltration process (see Table 30). The soluble BOD_5 removal of 89 percent for reverse osmosis alone was increased to 92 and 94 percent, respectively, by either the ultrafiltration pretreatment or the addition of activated carbon to the reverse osmosis feed. The advantageous use of activated carbon in the reverse osmosis feed, which had been pretreated by ultrafiltration, substantiates the previous conclusion that the high molecular weight materials of the ultrafiltration feed were utilizing or plugging all of the available adsorptive sites on the carbon.

Ferric chloride was added to the ultrafiltration feed in order to develop a floc and possibly increase membrane rejection and soluble BOD_5 removal. Ferric chloride actually worked in reverse and increased the transfer of soluble BOD_5 through the membrane. This loss of membrane rejection reduced the removal of the soluble BOD_5 components by 4 percent. Subsequent tests indicated that the ferric ion did not damage the membrane: after the module was washed with detergent, it gave salt (NaCl) rejection comparable to those for a new membrane.

SECTION VI

PILOT SCALE CLARIFIER STUDIES

GENERAL

The information from jar test studies of the effluents from the Mosinee and Combined Locks mills were used to prepare a scheme for the treatment of these mill effluents with a pilot-scale clarifier.

This clarifier had been developed at The Institute of Paper Chemistry under Project 3029, jointly sponsored by member companies of the Institute, the Environmental Technology Corporation (ENCOTEC) and the Upper Great Lakes Regional Commission (12). The stainless steel cone of the clarifier had a center height of 1.3 meters (4.25 feet) and a diameter of 1.1 meters (3.5 feet). The inlet was downflow through a 5.1-cm (2-inch) stainless steel pipe with an opening 40 cm (1.25 feet) from the bottom of the cone. It was equipped with an overflow weir which discharged into a 5.1-cm (2-inch) trough around the entire circumference of the top.

This section of the report covers the initial laboratory "shakedown" studies, the on-site studies at the mill in Mosinee, Wisconsin from January 15 to April 4, 1975, and the on-site studies at the mill in Combined Locks, Wisconsin from April 8 to June 26, 1975.

The problems encountered and some of the solutions developed for the correction of these problems are discussed. Data are given and discussed for the removal rates attained for suspended solids, total and soluble BOD₅ and color.

TRIALS WITH THE PILOT CLARIFIER UNDER LABORATORY CONDITIONS

Three thousand gallons of Mosinee clarifier influent were delivered by truck to the Institute early in January 1975 and were used to check the operation of the clarifier system (Figure 8) without the samplers or the submersible pump (P-1).

The system consisted of the following items and arrangement:

- 1. A tank (T-1) provided a constant supply of waste to be treated.
- 2. A feed control pump (P-2) maintained a constant feed rate.
- 3. A 35-liter (9-gallon) reactor tank (T-2) with a high speed mixer provided rapid mixing of the waste with the primary flocculating agent and permitted gravity flow to the balance of the stream.
- 4. The flow from the reactor tank was into an open pipe extending to the bottom of the flocculator tank (T-3). If polymer was to be added, it was introduced at the top of this open pipe,

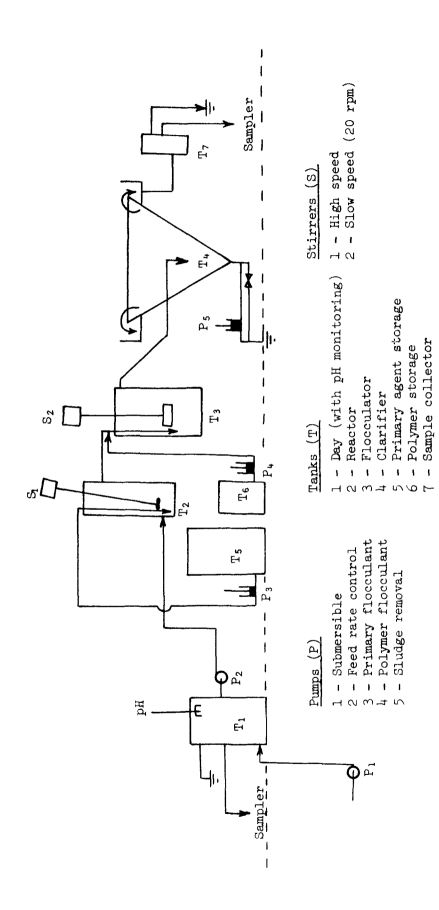


Figure θ . Schematic of pilot-scale clarifier installation.

into a quiescent zone approximately one-quarter of the distance down the pipe in order to provide adequate mixing without excessive turbulence or floc dispersion. In the center of this tank was mounted a slow speed (20 rpm = 1.6 fps peripheral velocity) paddle that mixed without excessive shear.

- 5. The overflow from the flocculator tank passed downflow into the clarifier (T-4), with the flow from the weir trough into a small sampling pot (T-7) and then to the sewer.
- 6. In these laboratory studies the sludge was removed from the bottom of the clarifier by gravity flow through a vented overflow pipe (not shown) that could be raised and lowered to control the rate of flow. Both gravity flow and a pumping system were used in the field trials and are discussed in the following part of this section.
- 7. The flocculating agents were stored in small tanks (T-5,6) and were metered into the system with tubing pumps (P-3,4).

We used this equipment to do a brief laboratory study of the effects of waste flow and sludge removal rates, at a constant level of flocculating agent(s), on the quality of the discharge produced. An excellent quality discharge (Table 31) appeared to be produced if the waste feed and the sludge discharge rate resulted in an overflow rate of 140-150 liters per square meter per day (400-425 gallons per square foot per day). The removal also appeared to be dependent upon the type of sludge blanket developed in the clarifier. Best results were obtained with the level at least five centimeters (2 inches) from the top to prevent floc carryover, but not less than one meter (3 feet) from the bottom; i.e., 60 cm (2 feet) above the waste inlet. This arrangement seemed to promote the development of larger floc particles and a filtration effect to markedly enhance the quality of the discharge. These parameters were, therefore, selected for the operation of the unit during the field trials.

EQUIPMENT AND OPERATION AT MOSINEE PAPER CORPORATION

The IPC system was to be operated in parallel with the two mill clarifiers and was, therefore, installed in the waste treatment plant in a concrete building. This building was immediately adjacent to the two commercial clarifiers and was unheated, except for heat from the incoming waste stream.

Due to the arrangement of the waste stream flow pattern, it was necessary to use a submersible pump (P-1 in Figure 8), mounted in the waste treatment trench immediately downstream from the bar and traveling screens, to lift the waste to floor level (2-2.5 meters or 6-8 feet) at a rate of 19-30 liters (5-8 gallons) per minute.

The flow from the submersible pump was directed into the bottom of a 190-liter (50-gallon) stainless steel tank (T-1) for supplying a centrifugal feed pump (P-2) with the excess overflowing to the sewer near the influent sampler. This tank also contained the pH recording equipment.

The feed pump (P-2) maintained a waste flow of 11.4 liters per minute (3 gpm) into the reactor tank (T-2) and the only other modification of the

equipment, over that described in the laboratory trials, was the installation of a pump in the sludge removal line during some of the trials.

TABLE 31. LABORATORY EVALUATION OF PILOT-SCALE CLARIFIER a

Over-	Sludge	Floccula	nt, mg/l		% Remova	al of	
${\tt flow}_{\tt b}$	removal,	Primary	Polymer	Suspended			OD_5
rate	% 	Fe ³⁺	н812.3	solids	Color	Total	Soluble
369	20	87	1.3	99	55	24	8
395	13	64	0.78	99	79	34	17
406	10	61	0.77	99		29	9
425	10	52	0.77	97	46	36	10
526	15	51	0.77	19	49	19	13
579	12	51	0.77	- 52	49	5	16
587	10	50	0.77	- 29	48	0	0
632	14	50	0.75	- 35	49	9	9

^aInitial (control) values: Suspended solids = 234 mg/l

Total $BOD_5 = 134 \text{ mg/l}$

Soluble BOD₅ = 114 mg/1

SAMPLING AT MOSINEE PAPER CORPORATION

During the first part of the mill study, influent and effluent samples were taken with two Model WM-5-24R refrigerated Sigmamotor samplers (Sigmamotor, Inc. of Middleport, New York) set to take individual samples hourly around the clock. In order to reduce the analytical load, these were composited into 12-hour samples daily.

It soon became apparent that the rapid changes in mill discharge (Table 32 and Figure 9) would not permit such an infrequent sampling scheme. A third sampler, a CVE refrigerated unit (Quality Control Equipment Company of Des Moines, Iowa) capable of taking small samples at short intervals (1-60 minutes) for compositing in a 4-liter (1-gallon) bottle, was placed in the influent stream. This unit was set to take 40-ml samples at 15-minute intervals and the 12-hour composites were compared with those from one of the Sigmamotor units taking 400-ml aliquots from the same influent.

Poor correlation was evident (Table 33) and a second trial was made. This was with four samplers in the following arrangement:

a. The two Sigmamotor samplers on the influent and effluent streams set at 30-minute intervals, taking 24 400-ml samples in 12 hours.

Color = 130 units.

Overflow in gallons/square foot/day based on surface area of clarifier.

- b. The CVE sampler on the influent line taking a 40-ml sample every 10 minutes.
- c. A polystaltic tubing pump (Buchler Instruments of Fort Lee, New Jersey) set to take 9 ml per minute continuously was used on the effluent line.

TABLE 32. ANALYSIS OF INDIVIDUAL SAMPLES TO ESTABLISH RANGE

		Mill	igrams/lite	er
		Suspended	Total	Soluble
No.	Нq	solids	BOD ₅	BOD ₅
1	10.12	210	241	203
2	10.07	628	206	178
3	9.57	494	166	138
4	9.63	145	155	138
5	9.11	85	129	116
6	9.48	144	135	114
7	8.86	265	143	140
8	9.40	152	163	140
9	9.19	97	148	140
10	9.30	76	127	117
11	9.78	94	138	135
12	9.60	115	132	124
13	9.59	267	148	118

Samples taken at hourly intervals (start 10 a.m. 1/16/75).

Analytical comparisons were excellent (Table 34) and indicated that the Sigmamotor samplers could be used to take representative samples if the 30-minute cycle was used.

In order to compensate for the "system holdup time" of 90 minutes, the samples were composited to obtain representative samples from the same 12-hour test period. This was readily accomplished by compositing the individually numbered bottles in groups so that those from the effluent were three bottles "behind" those for the influent.

Samples of the mill clarifier influent and effluent were taken at 7:30 a.m. daily by the mill staff and analyzed by both mill and IPC personnel.

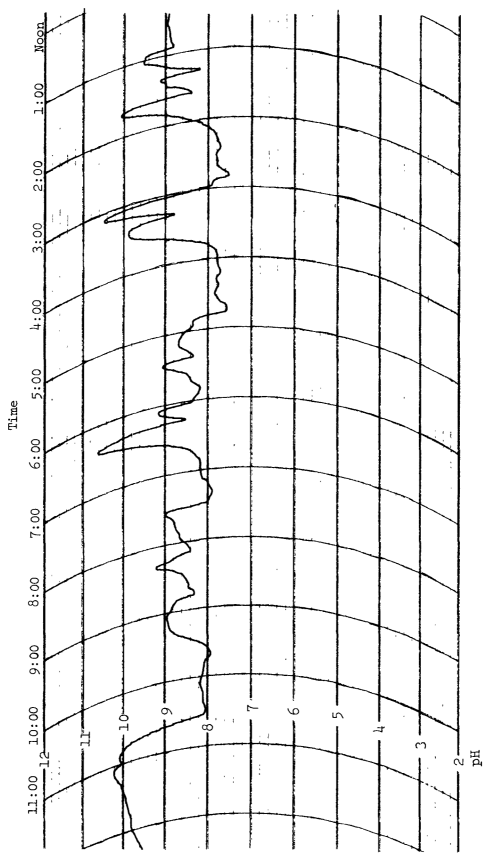


Figure 9. Variation in pH of Mosinee mill effluent entering pilot-scale clarifier.

TABLE 33. ANALYSIS OF SPECIAL AND REGULAR SAMPLES OF CLARIFIER INFLUENT, TRIAL NO. 1

					Mg/1	1			Pt	Pt-Co
	í d	H	Suspended solids	d solids	Total BODs	BOD 5	Soluble BOD ₅	e BOD ₅	00	color
	Special Regular	Regular	Special	Regular	Special	Regular	Special	Regular	Special Regular	Regular
	7.22 7.85	7.85	219	189	162	190	88	63	38	78
	7.82	8.75	182	136	177	544	109	138	115	130
	8.05	7.73	221	126	130	128	102	88	119	133
	8.40	8.51	210	199	124	158	108	117	170	204
	7.48	8.40	257	240	104	140	84	95	29	168
	7.35	7.61	213	169	86	142	9/	88	45	74
	7.35	7.11	213	198	120	88	76	98	09	82
ļ										

^aSpecial sample: A composite of 4 O-ml samples taken with a CVE sampler every 15 min for 12 hours; regular sample: a composite of 4 OO-ml samples taken with a Sigmamotor sampler every hour for 12 hours.

ANALYSIS OF SPECIAL AND REGULAR SAMPLES OF. CLARIFIER INFLUENT, TRIAL NO. $2^{\rm A}$ TABLE 34.

Samp	Sample No.		Hd		Total	Total BOD5, mg/1	/1	Soluble	Soluble BOD5, mg/1	g/1	Color (Color (Pt-Co units)	ts)
Special	Special Regular	Special	Special Regular	Mi11	Special	Regular	Mi11	Special	Regular	Mi 11	Special	Regular	Mi 11
AA-1	A-1	8.08	8.09	r C	197	180	701	162	148	1,0	268	273	17.3
AA- 2	A-2	7.41	7.22	0/./	174	173	100	120	124	143	95	145	<u>}</u>
BB-1	B-1	4.41	4.67		128	122		116	107	,	42	45	1
BB-2	B-2	3.69	3.61	t	6	98	ı	06	83	ı	65	52	1
AA-3	A-3	7.61	7.40	1	183	165	, 1	142	135	7	120	122	C
AA-4	A-4	7.19	7.20	7.33	140	139	0/1	110	104	011	50	89	2
BB-3	B-3	3,99	3.96		111	106		101	86	1	22	55	ı
BB-4	B-4	3.30	3.32		95	89		06	986		50	70	,

^aAll samples composited on a 12-hour basis.

 $^{\rm b}$ and B: A composite of 400-ml samples taken by Sigmamotor samplers at 30-minute intervals on influent and effluent. AA: A composite of 40-ml samples taken by a CVE sampler at 10-minute intervals on

influent.

BB: Continuous sample taken by a polystaltic pump at 9 ml/min from the effluent.

7:00 a.m. to 7:00 p.m. 7:00 p.m. 7:00 p.m. to 7:00 a.m. 7:30 a.m. composite. No. 1 & 3: No. 2 & 4: Mill:

For analysis all samples for BOD₅ were stored at 4°C (40°F) and the others were maintained at room temperature.

TESTING SCHEDULE AT MOSINEE CORPORATION

In order to establish a comparison between the mill and IPC clarifiers, the first trial was the operation of the pilot unit under conditions as near those of the mill as practical. Since we were not attempting to develop optimum clarification at high hydraulic loadings, but merely to study the increased removal of the soluble BOD5 from the waste stream while operating under different chemical treatment conditions, it was not necessary to show that we could exactly duplicate mill operation. Rather, it was more important to show that any increase in BOD₅ removal was due to the chemical additives and not to the clarifier design alone.

We developed a "base line" by operating the IPC unit with Hercofloc 812.3 at 0.75 mg/l and with a waste flow rate of 11.4 liters/min (3 gpm) in parallel with the two mill clarifiers. The Hercofloc addition duplicated mill treatment. If a 10 percent sludge withdrawal could be maintained, the pilot clarifier overflow would be 1.6 liters per sq cm per day (400 gallons per sq ft per day), approximately one-half that of the commercial unit.

The efficiency for removal of BOD₅ in the small unit was slightly less than for the mill units under the same conditions and time period. This indicated that the design of the IPC clarifier did not promote more removal and that any increases found in later trials using chemical additive would be directly attributable to the chemical treatment.

Following the development of the base line, other chemical treatment schemes based on the laboratory studies were tested:

- A. Base line using 0.75 mg/l Hercofloc 812.3.
- B. Alum at 200 mg/l plus 0.75 mg/l Hercofloc 812.3.
- C. Ferric chloride at (100 mg/l as Fe³⁺) plus 0.75 mg/l Hercofloc
- D. Ferric sulfate (50-100 mg/l Fe³⁺) without the polymer.
- E. A second base-line study with the Hercofloc.
- F. Sulfuric acid for pH adjustment only.
- G. Sulfuric acid for pH adjustment plus the Hercofloc. H. Ferrous sulfate (100 mg/1 Fe²⁺) plus 0.75 mg/l Hercofloc.

The original plan was to have tests of two weeks' duration. Unfortunately, the several periods of problem and solution development we encountered early in the study delayed the program and resulted in shortened test periods (for F, G, and H, above) near the end of the study. Also, throughout almost the entire trial period, the Mosinee mill was on a reduced work week (4 days), which limited the operation of the small clarifier to the period between 9:00 a.m. Tuesday and 4:00 a.m. Saturday, thereby limiting the number of samples per week.

RESULTS AND DISCUSSION OF WORK AT MOSINEE

In our first attempt to establish the base line, we immediately encountered three problems:

The first problem related to the submersible, centrifugal sump pump installed in the waste trench. It had a small 2.5-cm (l-inch) inlet hole in the baseplate and due to the high fluid velocity in the waste trench, an "aspirator" effect developed across the inlet when the pump was vertically mounted, and we were unable to pump sufficient waste. Mounting the pump at a 90° angle pointed downstream increased the aspiration effect, while mounting it pointed upstream resulted in rapid plugging of the inlet hole with pitch and debris not removed by the bar or traveling screens. This problem also extended to the centrifugal pump used for feed control.

The second problem was due to the high suspended solids content of the Mosinee waste. A centrifugal pump was the only available pump that was considered suitable for control of the flow at the ll.4 liters per minute (3 gpm) rate. A one-half inch Eastern centrifugal pump was, therefore, used to control the rate of waste fed to the clarifier unit, and at first the rate was controlled with a 1/2-inch valve on the pressure side of the pump. This rapidly plugged with pitch and fibers and resulted in erratic flow patterns.

The third problem developed in the gravity sludge removal system. Settling of the high suspended solids content rapidly overwhelmed the available space in the clarifier and resulted in solids carryover, even in the absence of chemical agents, other than the Hercofloc 812.3.

In order to reduce these problems:

- l. A conical screen (20 mesh), 18 inches long by 7 inches in diameter, was installed on the submersible pump. When installed with the apex of the cone mounted upstream, the pumping action was excellent and the cone was somewhat self-cleaning. When mounted with downstream orientation, which should increase the self-cleaning aspect of the cone shape, the flow was reduced below acceptable levels. Periodic cleaning, with high pressure water, was required to maintain an adequate supply to the system.
- 2. The centrifugal feed pump was fitted with a smooth-bore orifice plate for flow control and with periodic cleaning this provided better flow control.
- 3. A multihead tubing pump (Brosites) was installed in place of the gravity system for sludge removal. In the last four trials a throttled centrifugal pump was tried in this position and found to work the best of any of the schemes attempted.

Although none of these worked perfectly and problems were encountered from time to time, failure of the system could be minimized with careful cleaning and operation procedures.

Data for the trials with the various flocculating agents and for the base lines are summarized in Table 35. The more detailed analysis of the individual samples are in Appendix B (Figures B-1 to B-10, Tables B-1 to B-5). These data were first used to establish confidence limits, and the following summary was developed.

- 1. Both the initial and final trials (A and E of Table 35) with 0.75 mg/l Hercofloc 812.3 indicated, as previously noted, that the IPC unit was equal to or slightly less efficient than the mill clarifiers in removing suspended solids, soluble BOD₅ and color.
- 2. The reason for the marked difference noted in the <u>total</u> BOD₅ reductions for the mill clarifier during the second trial is unknown; throughout the study period there had been a slow, progressive increase in the efficiency of the mill clarifier in removing BOD₅; i.e., from 14 to 35 percent between Trials A and H. This was not readily traceable to changes in the concentration of BOD₅ going to the clarifiers during this period, since these were on the average 161, 120, 138, 181, 161, 211, 114 and 178 mg/liter for Trials A to H, respectively.
- 3. Alum with polymer and without pH adjustment, as predicted by the laboratory trials, did not increase the removal of the BOD_5 components but did markedly increase the efficiency of removal of suspended solids and color.
- 4. The ferric ion, both as chloride and sulfate salts, was an excellent flocculating agent for removing BOD_5 , the chloride form being more effective than the sulfate for suspended solids. The chloride could have been better because the final pH was nearer to the optimum for the Fe³⁺ produced flocs (pH 4-5).
- 5. Ferrous ions, at the level of 100 mg Fe²⁺/l, appeared to reduce the efficiency of the system even with the addition of polymer. While this level of ion was below that which would be theoretically required by the Schulze-Hardy rule, which states that the concentration of the counterion required to collapse the double layer and produce rapid flocculation is inversely proportional to the 6th power of the valence, it was the highest level practical during our trials due to the low solubility of the commercial grade ferrous sulfate. Based on the effectiveness of the ferric ion at 100 mg/l, the level of ferrous ion theoretically required would have been approximately 1100 mg/liter.
- 6. The use of sulfuric acid to decrease the pH was one process under consideration by the mill for use in the large clarifiers. A trial was, therefore, designed to test this procedure for the removal of some of the components from the waste stream in the small system; but due to the mechanical difficulties we had encountered earlier in the trial period, only a short period was available near the end of the study. Although the data were insufficient to provide reliable comparisons with mill operation, laboratory studies have shown that pH reduction to below 4.5 resulted in marked reduction in suspended solids and total BODs, with little effect on the soluble BODs level in the effluent.

TABLE 35. VARIOUS TREATMENTS OF MOSINEE MILL EFFLUENT IN PILOT-SCALE AND MILL CLARIFIERS

These trials, with ferric, ferrous, and aluminum ions, and especially with the iron salts, have shown that an increase in clarifier efficiency (12 percent for total BOD₅, 15 percent for soluble BOD₅ and up to 66 percent for color) could be attained with the Mosinee waste. The sulfate form might be preferable to the chloride, particularly if the sludge were to be burned, because it is less corrosive.

One of the principal practical observations we made in the field study at this mill was on the effect of strong digester room spent liquors and washes on clarifier performance. Visual, analytical, and other evidence showed that heavy slugs and spills of these wastes — some of which lasted several hours — adversely affected clarification. A single program of grab sample collection was undertaken to help indicate the sources of strong process waters. The analytical data from the study of that one set of grab samples are summarized in Table 27, previously described, and showed the relative strength of the various soluble materials in these samples. The effects of these streams will be discussed in greater detail in Section VII.

EQUIPMENT AND OPERATION AT COMBINED LOCKS, APPLETON PAPERS DIVISION OF NCR

The clarifier equipment that had been used in the Mosinee field trials was cleaned and transferred to Combined Locks on April 7, 1975. Since it was again to be operated in parallel with the mill system, it was placed in the treatment plant building which was immediately adjacent to the two mill clarifiers. A tee was installed in the main influent line to these two clarifiers and this stream was fed into the 50-gallon day tank (T-1 in Figure 8). This eliminated the need for the submersible pump. At Mosinee we had encountered some difficulty with uneven flow over the weir of the small clarifier. Although the unit had three leveling legs, the top rim was not level around the entire periphery. Spot overflow occurred; the floc subsequently "streamed" at these points and efficiency decreased.

An adjustable leveling device for the top of the clarifier at Combined Locks was made of small strips of slotted plastic installed along the entire overflow edge. Carefully manipulated (up and down), these could be used to establish an overflow around the entire circumference. This helped to maintain a smooth top on the sludge blanket and virtually eliminated spot overflow except when the strips were disturbed during the weekly cleanup.

Two other basic changes were made for operation at the Combined Locks mill to provide better control. The high fiber content of this waste stream overwhelmed the sludge disposal system on the IPC clarifier. In order to reduce the fiber content and still retain in the influent the fine particles required for good floc formation and BOD₅ removal, we installed a side hill screen with 60-mesh wire over the top of the day tank. Later, when we encountered difficulties in maintaining a clean surface in the base-line study (A), we changed this to a Hydrasieve (C. E. Bauer, Springfield, Ohio) with 0.010 inch slots (Figure 10).

A centrifugal pump with electrical speed control was used to control the sludge level in the clarifier and the hose was clamped on the outlet side.

This was changed to run at low speed with the hose clamp removed, and the time of operation was controlled with a timed ON/OFF switch (Trial D). This permitted relatively close control of the sludge blanket level that had varied both with the "quality" of the waste fed to the system as well as with the type of flocculating agent.



Figure 10. Bauer Hydrasieve used to remove the longer fibers from the Combined Locks mill effluent

Since the level of the sludge blanket in the clarifier was apparently very important to the efficiency of the system, thought should be given to provision of a sensing type of control system for any future work with this unit.

SAMPLING AT COMBINED LOCKS

The two Sigmamotor samplers were used on the influent and effluent streams. Comparisons were again made with these two samplers, the CVE sampler and the polystaltic pump. The Sigmamotor samplers were set to take 400ml samples every 30 minutes, the CVE sampler a 50-ml influent sample every 8 minutes, and the tubing pump a continuous 9 ml/min sample of the effluent. Samples from each were composited every 12 hours.

Data in Table 36 indicate that representative samples were taken by the Sigmamotor samplers set at the 30-minute sampling frequency. To provide for the "system hold-up" we used the same "lag period" in compositing that we used at Mosinee. Samples of the mill clarifier effluent were taken by the mill staff every morning and were analyzed at the IPC laboratories for comparison with those from the small unit. The mill had no provisions for taking 24-hour composites of the mill clarifier influent, so the influent values for the IPC clarifier for the same test period were averaged for comparison of percentage removal values for total and soluble BOD5 and color. Since the Hydrasieve removed considerable quantities of fiber from the influent processed by the IPC unit, the removal of suspended solids could not be compared with that of the mill system.

TESTING SCHEDULE AT COMBINED LOCKS

The Combined Locks clarifier operates without chemical additives and depends upon gravity sedimentation for clarification. The "base-line" trial (Trial A) of the IPC unit was, therefore, made without chemicals. This was followed by trials:

- B. Ferric chloride at 75 mg/l Fe3+.
- C. Ferric chloride at 75 mg/l Fe³⁺ with 0.75 mg/l Nalco 73C32. D. Ferric sulfate at 100 mg/l Fe³⁺.
- E. Alum at 300 mg/l.
- F. Alum at 300 mg/l with 0.75 mg/l Hercofloc 812.3.
- A second base-line study with no chemical additives.

Because the IPC system was not automatically controlled and because clarifier influent quality changed rapidly on Saturdays and Sundays, we usually limited our trials to 8:00 a.m. Monday through 7:00 a.m. Saturday; several trials, however, extended through the weekend.

RESULTS AND DISCUSSION

The Hydrasieve, cleaned frequently, provided good influent for the IPC clarifier. The basic problems at this site were with plugging of the influent line between the mill main line and the 50-gallon per day tank and with the removal of sludge from the small clarifier to maintain proper sludge-

TABLE 36. ANALYSIS OF SPECIAL AND REGULAR SAMPLES OF THE CLARIFIER INFLUENT OF THE COMBINED LOCKS MILL^B

	Sample no.	e no.	Hq		Suspended mg/l	Suspended solids, mg/l	Total BODs, mg/l	0s, mg/l	Soluble BODs, mg/l)D ₅ , mg/1	Color	or
Day	Special ^b	Regular	Special ^b	Regular ^c	Special ^b	Regular ^c	Special ^b	Regular ^c	Specialb	Regular ^c	Special ^b	Regular
2	DD-1 DD-2	D-1 D-2	8.59 7.68	8.50 7.70	62 77	7.1 7.4	291 304	273 296	270 250	275 251	170 237	170 237
m	DD-1 DD-2	D-1 D-2	8.00	7.89	333 220	324 201	382 370	351 344	752 757	264 247	228 203	216 222
4	DD-1 DD-2	D-1 D-2	7.78	7.70	134 191	104 182	275 280	285 256	211 224	216 210	285 240	287 258
72	DD-1 DD-2	D-1 D-2	7.80	7.89 7.71	68 105	60 97	224 264	227 256	198 221	202 201	200	192
80	DD-1 DD-2	D-1 D-2	8.01	8.68 7.90	07	14.7 3.7	201	228 197	170	196 184	1 1	155 150
0	DD-1 DD-2	D-1 D-2	8.31 8.03	8.20	59 142	51 41	250 264	250 286	214 259	218 228		183 170
10	DD-1 DD-2	4 4 4 8	7.65	7.82	19 11	71 61	148 214	162 208	138 186	140 176	; ;	105 150
11	DD-1 DD-2	D-1 D-2	7.71 7.82	7.62	70 35	66 37	234 272	240 282	216 249	211 238	11	21 <i>7</i> 302
12	DD-1 DD-2	D-1 D-2	7.62	7.50	19	19 18	128 166	140 162	125 170	128 154	85 87	75 83
13	DD-1 DD-2	D-1 D-2	7.73	7.65	32	44 31	264 284	270 286	256 255	254 271	225 210	200 205
14	DD-1	D-1	7.63	7.58	65	38	252	243	235	250	241	255
8	,											

aAll samples were composited on a 12-hour basis. bSpecial samples: 50 ml taken by a CVE vacuum sampler at 8-min intervals. Regular samples: 400 ml taken by a Sigmamotor sampler at 30-min intervals.

blanket levels. The first remained a recurring problem throughout the study. The sludge blanket was fairly adequately maintained with the installation of the time-controlled centrifugal pump.

Data from these trials (Tables C-1 to C-7 and Figures C-1 to C-14 in Appendix C) were checked for validity (mathematical confidence limits) on the IPC computer. Averaged data for the seven trials are given in Table 37 and are summarized as follows:

- 1. The first base line (Trial A) showed that the IPC clarifier removed slightly more total BOD₅ than the mill unit did, but removed equal amounts of soluble BOD₅ and no color. In the second base line (Trial G) at the end of the series the units operated at equal efficiency.
- 2. Ferric ion, either as the chloride or sulfate salt (Trials B and D) provided excellent removal for total and soluble BOD₅ as well as color.
- 3. The addition of the polymer Nalco 73°32 (Trial C) to the ferric chloride slightly reduced the efficiency for total and soluble BOD₅ removal but had no effect on the color removal.
- 4. Alum, as predicted by the laboratory jar tests, did not appear as effective as the ferric ion in removing BOD₅ but was equivalent to it in reducing color.
- 5. The addition of the Hercofloc 812.3 polymer to the 300 mg/1 alum markedly increased the removal of total BOD_5 to well above the level achieved with the other flocculating agents but had no effect on the reduction of the soluble BOD_5 .

The high "negative" soluble BOD₅ removal in the two IPC base-line trials (Appendix C-1 and C-7), as well as throughout the test period for the mill clarifier, could not be explained. This finding, however, did correlate with the darker color of the sludge from the mill clarifiers (lighter fibrous material was in the influent waste stream) and might indicate development of both color and soluble BOD₅ from the "particulate matter" during clarification.

Spent liquor entering the influent system through spills or deliberate discharges had the same deleterious effect in this study at Combined Locks as it did at Mosinee. This, according to the Combined Locks mill staff, has been one of the causes of clarifier upsets that have occasionally plagued their operation. Operation of the pulp and paper mill under strict control of spills and similar discharges (i.e., good housekeeping) has minimized these upsets.

In our laboratories we studied the effect of the various sewer discharges on clarification, and the data developed in these studies are discussed in detail in Section VII.

TABLE 37. CLARIFIER STUDY — COMBINED LOCKS PAPER CORPORATION AVERAGE EFFLUENT QUALITY AND REMOVAL VALUES FOR IPC AND MILL CLARIFIERS

		Suspended	Effluent c	uality D ₅			Removal, %	
		solids,	Total,	Soluble,	Color		OD ₅	
Cla	rifier	mg/l	mg/l	mg/l	units	Total	Soluble	Color
Α.	Both un	nits without	chemical	flocculati	ng agents	:		
	IPC Mill	46 48	232 254	190 188	205 227		8.6 8.3	-20.6 -33.5
В.	IPC uni	it with ferr ent:	ic chlorid	le (75 mg/l	Fe ³⁺), an	d mill u	sing no che	emical
	IPC Mill	45 74	169 214	147 187	112 207	40.1 24.1	25.8 5.6	32.5 -24.7
С.		it with ferr ll using no			Fe ³⁺) +	Nalco 73	C32 (0.75 m	ng/l),
	IPC Mill	66 46	196 214	168 209	92 199	34.4 28.4	20.4	31.8 -47.4
D.	IPC unt	it with ferr ent:	ic sulfate	e (100 mg/l	Fe ³⁺), an	d mill u	sing no che	emical
	IPC Mill	37 50	136 166	129 152	77 130	41.9 29.0	25.4 12.1	
E.	IPC un	it with alum	(300 mg/]	l), and mill	using no	chemica	l treatment	::
	IPC Mill	60 54	205 240	197 216	123 241	_	13.2 4.8	31.7 -33.9
F.		it with alum no chemical			loc 812.3	(0.75 m	g/1), and m	ill
	IPC Mill	85 56	237 262	208 212	129 229	50.9 30.9	13.3 11.7	30.3 -23.8
G.	Both unonly):	nits without	chemical	flocculati	ng agents	(gravit	y sedimenta	ation
	IPC Mill	50 52	239 236	214 208	252 254	27.8 28.7	-1.4 1.4	-26.6 -27.6

^aMill did not sample influent; values are based on average of IPC influent.

SECTION VII

JAR TESTS WITH INDIVIDUAL SEWER DISCHARGES

GENERAL

In Section V of this report, the BOD₅ sources making up the total discharge of the mill to the clarifier were listed and briefly discussed. In the field studies (Section VI) we had noted a marked decrease in the efficiencies of both the mill and IPC clarifiers when larger than usual amounts of spent liquor were in the waste stream being processed.

In this section we detail and discuss the studies made on the efficiency of the soluble BODs removals realized when individual sewer streams were treated. These individual sewer effluents were also used to make a "synthetic" mill effluent by compositing each on the basis of its volume to the overall volume of the total mill discharge.

Using these same individual streams, we prepared various composites, omitting one stream at a time, to determine the effect the omission had on the overall efficiency of BOD₅ removal by clarification.

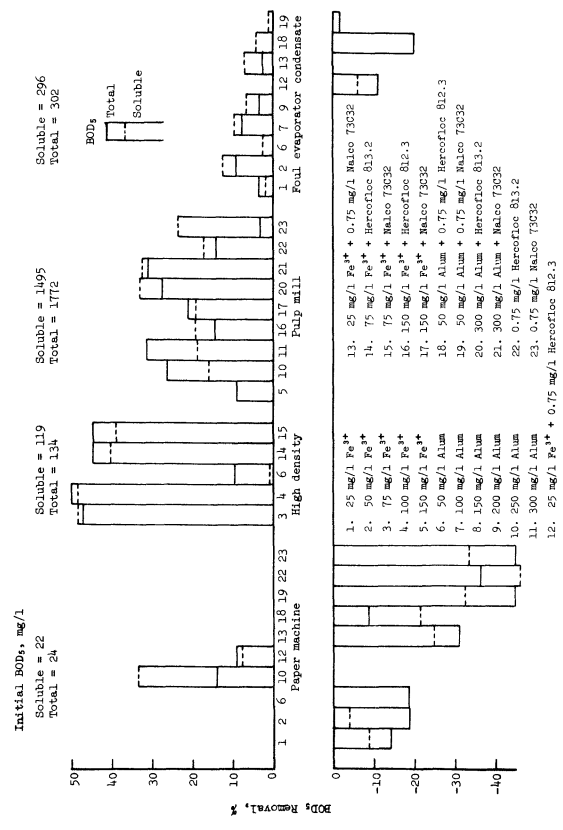
Ferric chloride (25, 50, 75, 100, and 150 mg/l Fe³⁺), alum (50, 100, 150, 200, 250, and 300 mg/l) and combinations of these two primaries with 0.75 mg/l of either Hercofloc 812.3 or Nalco 73C32 were used with each stream and composite.

Values for total BOD_5 and, when time permitted, soluble BOD_5 for the treated samples were compared with untreated (settled) control samples for the determination of BOD_5 removal.

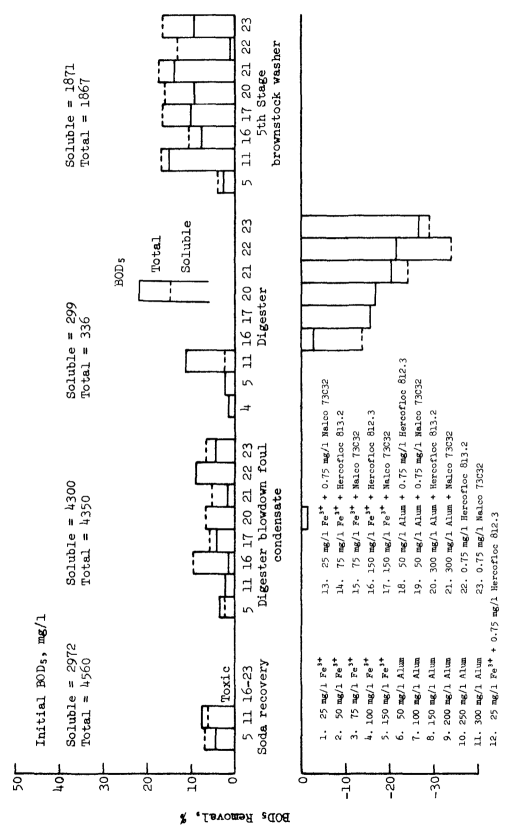
TESTS ON MOSINEE INDIVIDUAL SEWER SAMPLES

Some of the mill discharge streams, especially the "high density," pulp mill and paper machine sewers, could be effectively treated with the proper concentration of the primary flocculants (Figures 11 and 12, Legends 1-11) or primary flocculant-polymer combinations (Legends 12-23). These bar graphs display that in others, such as the foul evaporator condensate, soda recovery wastes, and digester room wastes, the BOD₅ level was not substantially affected by any of the flocculating agents. Those bars shown below the zero line in the lower section of the figures indicate negative removal of BOD, probably due to prolonged holding time. These waste flows should be treated by routes other than clarification (i.e., steam stripping of the condensates).

In our earlier work with the individual streams we found that toxic materials were in some samples, and they interferred with the BOD_5 test (Table 38). Therefore, we could not make even good estimates of the concentration



Jar tests on wastes from individual sewers in the Mosinee mill (paper machine, high density, pulp mill, and foul evaporator condensate). Figure 11.



Jar tests on wastes from individual sewers in the Mosinee mill (continued) (soda recovery, digester blowdown, foul condensate, digester, and 5th stage brownstock water). Figure 12.

TABLE 38. BOD OF INDIVIDUAL EFFLUENT STREAMS AND TOTAL MILL EFFLUENT OF MOSINEE MILL

Source	Dilution	mg/l
Digester sewer	120 600	637 633
Pulp mill sewer	150 300 600 1500	1210 1490 1629 1620
5th Stage brownstock washer	300 600 1000 2000	940 1089 1353 1340
High density sewer	30 100 150	141 174 172
Foul evaporator condensate	120 300 600	570 762 751
Digester blowdown foul condensate	300 600 750	1023 1325 1300
Soda recovery sewer	600 1500	4236 4188
Total mill effluent	60 150	271 280

of BOD₅ in these streams without careful dilution techniques. Excessive amounts of these same toxic agents would also interfere with the operation of a biological secondary treatment system and for this reason, if for none other, their presence in the total effluent should be held to a minimum or completely eliminated. Discharges into individual mill sewers were used to prepare a "synthetic" total mill effluent in the following volume relationships:

	Million gal/day	% of Total
Paper machine	10	77.9
High density	1.5	11.7
Pulp mill	0.6	4.6
Foul evaporator condensate	0.3	2.3
Soda recovery	0.28	2.2

Digester blowdown		0.07	0.6
Digester		0.05	0.4
5th Stage brownstock washer		0.035	0.3
	Total	12.835	100.0

To find the conditions for maximal BOD_5 reduction when individual wastes were omitted from the system, we made combinations of seven of these eight wastes and treated them also in jar tests. Data in Figure 13 indicate that higher levels of BOD_5 removal could be achieved with less chemical if certain individual discharges were eliminated from the flow. That is:

- 1. The "total" effluent required 250 mg/l of alum for 17 percent reduction or 75 mg/l Fe $^{3+}$ for 42 percent reduction in BOD5.
- 2. With the soda recovery sewer flow eliminated, a 250 mg/l alum level resulted in 26 percent reduction and the Fe³⁺ at 50 mg/l resulted in a 51 percent removal of BOD₅.
- 3. The absence of the pulp mill sewer from the total flow, with all other flows present, permitted a 60 and 63 percent reduction in the BOD_5 concentration in the waste stream for 100 mg/l alum and 50 mg/l Fe³⁺, respectively.

When 1 percent v/v of the digester room waste was added to a mixture of wastes having relatively good clarification characteristics (paper machine and high density wastes), higher concentrations of flocculating agent were needed to produce an effluent of adequate discharge quality (Figure 14).

Attempts to calculate BOD_5 balances for these various discharge combinations, and to correlate these with the actual BOD_5 values obtained, was apparently hindered by some interaction(s) between these wastes. This could have been due to changes in the toxicity of the mixtures for the microorganisms in the BOD_5 test.

The difficulty in calculating material balances for the total or soluble BOD₅ of the individual streams and composites is due to the fact that all of the initial BOD₅ values for these samples were on "settled" jar samples. Settling removes BOD₅ and the removed BOD could not be accounted for in the calculations of BOD₅ balances. In these samples, the interaction of the adsorbing character of fibers and particles with the dispersing character of the various mixtures could markedly affect BOD₅ removal during the settling step, thereby modifying the "initial" BOD₅ of the mixture. The importance of this interaction in the overall flocculation-clarification process should be investigated further in any attempt to improve the process.

TESTS ON INDIVIDUAL STREAMS FROM COMBINED LOCKS MILL

The five streams making up the bulk of the flow to the clarifiers at Combined Locks — from the main, "CM," wet room, Number 5 and Number 6 sewers — were tested individually for the removal of BOD₅ in jar tests with ferric chloride, alum and combinations of these with either Hercofloc 812.3 or Nalco 73C32. We also prepared a composite "total" mill effluent and composites lacking individual sewer outfalls, as we did with the Mosinee wastes.

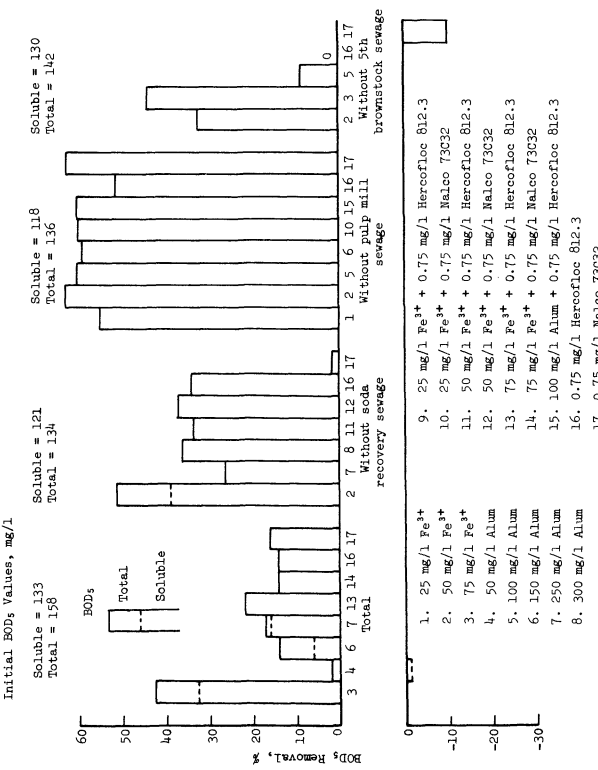


Figure 13. Jar tests on special composites of Mosinee individual waste streams.

As predicted by our earlier work, those streams having low spent liquor content, i.e., Number 5 and main sewers, had good supernatant quality (Figure 15). The BOD₅ removal was good with relatively low concentrations of Fe³⁺ or alum (Figure 16).

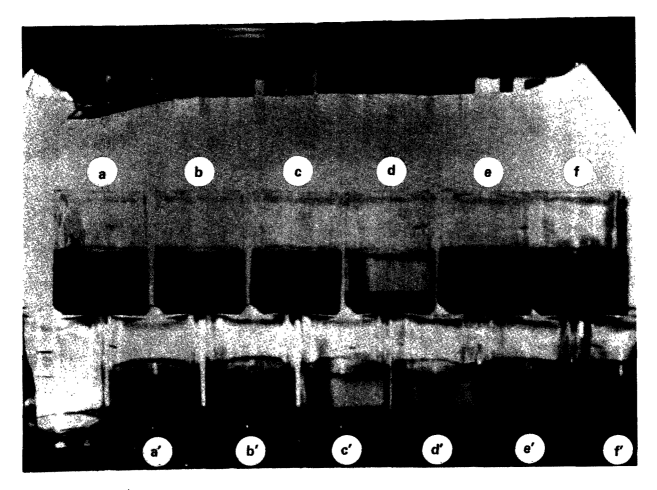


Figure 14. Jar tests show mixed effluents with addition of, left to right, 25, 50, 75, 100, and 150 mg/l Fe³⁺; the control, no addition, is jar f. A Mosinee effluent was used composed of: (bottom row) paper machine and high density sewer effluent, and (top row) the above with 1 percent v/v digester room effluent.

The chemical concentrations of ferric chloride and alum required to effectively remove both suspended solids (for high supernatant clarity) and BOD_5 was higher for the "total" effluent than for the various composite streams (Figure 17). As the individual streams were removed from the system, the quantity of flocculants was reduced, especially for those mixtures without the "CM," wet room or Number 6 sewers. The Number 5 sewer stream had good removal characteristics, with low chemical requirements, as an individual stream, and omitting it from the mixture did not appear to enhance the BOD_5 removal.

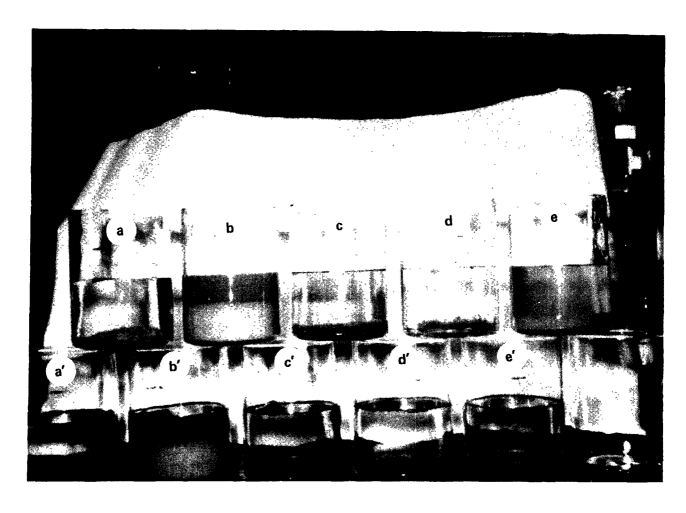


Figure 15. Jar tests with Combined Locks individual sewer effluents, a-e (main, "CM," wet room, machine No. 5 and machine No. 6). Top row: with 25 mg/l Fe³⁺; bottom row: with 20 mg/l alum. Control wastes did not settle without additives and are not shown.

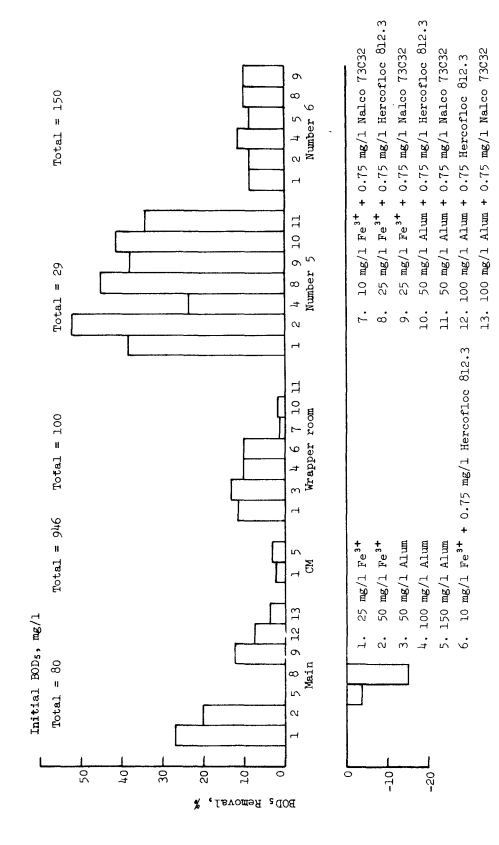


Figure 16. Jar tests on wastes from individual sewers in the Combined Locks mill.

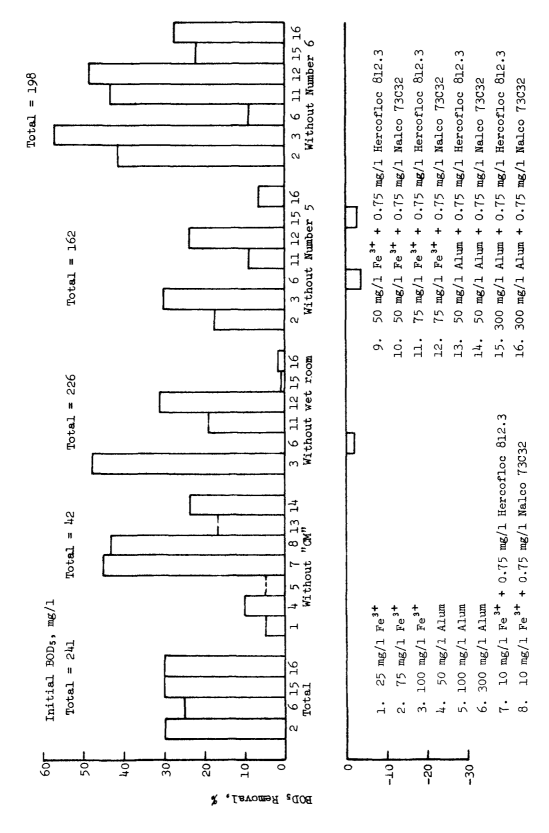


Figure 17. Jar tests on special composites of Combined Locks individual waste streams.

SECTION VIII

DISCUSSION OF THE MECHANISMS OF SOLUBLE BOD5 REMOVAL

The studies with jar and pilot-scale clarifiers have readily confirmed that substantial quantities (as much as 20 to 50 percent) of the soluble BOD₅ are removed in clarifiers by coagulation aids and that the removal can be a significant factor in the treatment of pulp and paper mill waste waters.

Our data show that, at least to some extent, the reactions leading to improved BOD₅ removal can be optimized in well operated and controlled clarifier systems. Several studies were made in an attempt to better define the mechanism for reduction of soluble BOD₅ during the treatment of process waters in primary clarifiers.

It had been evident from the beginning of the study that some of the soluble BOD₅ could be removed from the waste stream by several processes (membrane, ion-exchange) and that no one mechanism could be credited for the entire effect.

GEL CHROMATOGRAPHY STUDIES

The components responsible for the soluble BOD₅ in clarifiers can be divided into two classes, wood-derived chemicals and additives. The wood derived chemicals include degraded cellulose, hemicellulose, hemicellulose degradation products (reducing sugars, acetic acid, methanol and uronic acids), extractives and in small, perhaps minor amounts, metabolites common to living tissues (adipic acid, oxalacetic acid, etc.). Additives used in papermaking include starches, polysaccharides, gums, latexes, resins, dyes, etc. Some mills will also have organic solvents such as methanol, acetone, etc., which are biodegradable. The organic solutes that pass the 0.45-µm filter probably comprise colloids, low molecular weight polymers, oligomers, monomers and simple organic compounds.

To better understand the removal of soluble BOD₅ by coagulation, we treated the effluent from a chemimechanical mill (Combined Locks mill) with the various flocculating agents. Using gel chromatography we measured molecular size distribution before and after flocculation and clarification.

Samples were filtered on a 0.45-µm filter prior to chromatography. The gel chromatography was carried out on Sephadex G-50, a gel which excludes (allows no equilibration with the gel pores) molecular weights 10,000 and above and includes (allows complete equilibration with the gel pores) molecular weights of 500 and below. Molecular weights between 500 and 10,000 are fractionated according to the number of pores with which the molecules equilibrate.

Figure 18 shows the TOC (total organic carbon) of 2 ml of a tenfold concentrate as it elutes from a 1 x 55-cm column. The second curve shows that after flocculation with 70 mg/l ferric chloride not only are large molecular weight residues removed, but also low molecular weight molecules. The curves in Figure 19 were obtained when a similar sample (1-ml sample of fivefold concentrate) was fractionated on Sephadex LH 20 (mol. wt. 5000 to 200) before and after coagulating with 10 mg/l of Nalco 73C32, a cationic polyelectrolyte and analyzed for aldoses by the phenol-sulfuric acid test (8). The low molecular weight peak almost disappeared.

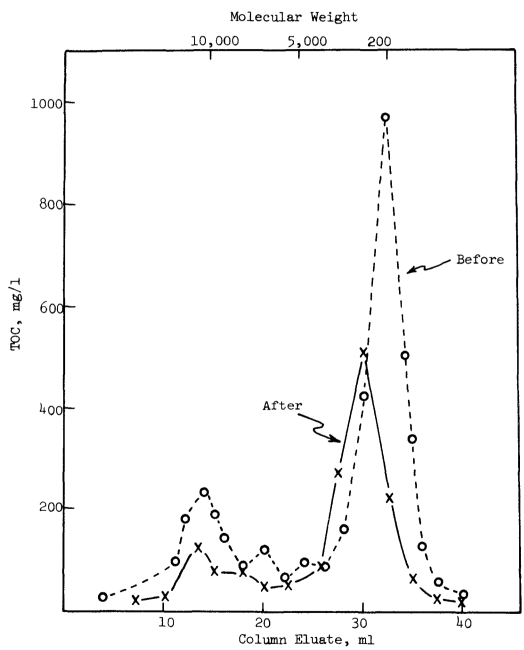


Figure 18. TOC and molecular weight distribution of gel-chromatographed chemimechanical mill effluent before and after coagulation with 70 mg/l ${\rm Fe}^{3+}$ as ferric chloride.

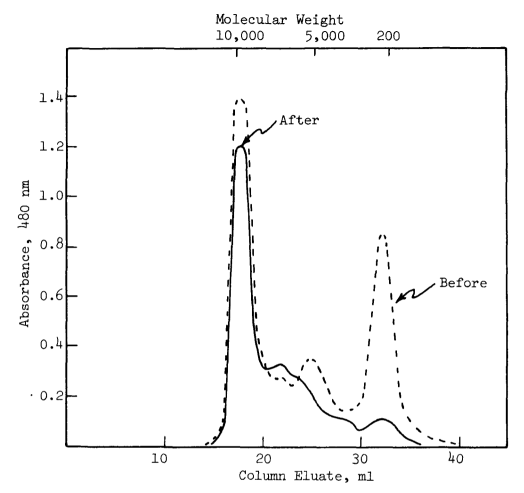


Figure 19. Phenol sulfuric acid test for aldoses and molecular weight determination of gel-chromatographed chemimechanical mill effluent before and after coagulation with 10 mg/l Nalco 73C32. One-ml samples of eluate were analyzed.

The phenol-sulfuric acid test measures aldoses and polymers of aldoses when borate interference is observed. The solutions analyzed in this report showed enough borate sensitivity to indicate the test was responding primarily to aldoses and polymers of aldoses and not to noncarbohydrate aldehydes.

When a sample fractionated on Sephadex G-50 was divided into three fractions corresponding to high, medium and low molecular weights and analyzed for TOC, aldose and BOD₅, the results summarized in Table 39 were obtained. The percentage change for the soluble BOD₅ was not as great as that observed for the aldoses or TOC. Apparently not only low molecular weight biodegradable materials were removed but also low molecular weight nonbiodegradable organics.

Das and Lomas explored the different mechanisms of the flocculation of cellulose fines with the cationic polymer polyethylenimine (2). They were

able to show that bridging (the adsorption of two sections of a polymer to two colloids) was a significant part of the cellulose fines flocculation. Such a mechanism could also occur with the other polyelectrolytes and with ferric chloride which hydrolyzes and polymerizes to a polyelectrolyte (14). The mechanisms for the coagulation of cellulose fines involved collapse of their double layer followed by adsorption and bridging. Lower molecular weight residues were generally difficultly adsorbed. Their participation in coagulation probably depends on salt formation and chelation. Such salt formation by hydrolyzed iron can be demonstrated with gel chromatography. Figure 20 shows the elution curve for a mixture of ferrous ammonium sulfate and sodium gluconate. The complexes absorb ultraviolet light and are, therefore, detectable with the UV absorptiometer at 280 nm. Iron values were obtained by the o-phenanthroline test. Because the curve shows that at a pH of 2.0, most of the iron elutes from Sephadex LH-20 and G-15 (fractionation range 150 to 10,000) in the highest molecular weight fraction, the complexes probably have molecular weights of 10,000 and above. Figure 21 shows the distribution of the complexes when the pH of the solution was 7.0. At higher pH more of the iron was in the lower molecular weight fractions.

TABLE 39. REMOVAL OF DIFFERENT MOLECULAR WEIGHT FRACTIONS OF ORGANIC RESIDUES BY FLOCCULATION

		Sam	ole ^a	
Effluent treatment	UNF	HMW	MMW	LMW
Phenol-sulfuric acid test; total absorbance, 480 nm				
Filtered Flocced ^b and filtered Reduction, %		7.5 3.0 59.9	3.9 2.4 38.6	5.2 2.5 51.9
TOC, mg/l				
Filtered Flocced and filtered Reduction, %	3424 2150 43.9	774 320 58.6	457 272 40.5	2301 1387 39.7
BOD ₅ , mg/l				
Filtered Flocced and filtered Reduction, %	3840 3080 19.8	552 379 31.3	487 368 24.4	2689 2131 20.7

^aUNF, unfractionated; HMW, high molecular weight (10,000 and above); MMW, middle molecular weight (500-10,000); LMW, low molecular weight (500 and below).

Flocced with ferric chloride, 70 mg/l Fe.

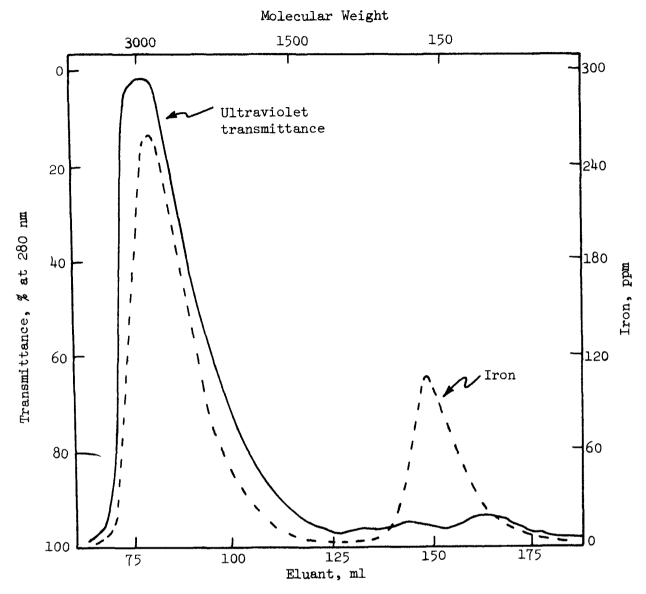


Figure 20. Ultraviolet light transmittance profile and molecular weight distribution of a gel-chromatographed solution of sodium gluconate and ferrous ammonium sulfate, pH 2.

If these biodegradable low molecular weight residues which participate in the coagulation are carboxylic acids, then the mechanism for their removal could be the complexing and/or chelating of the Fe³⁺ of the ferric chloride. As the ferric chloride hydrolyzes, more and more insoluble oligomeric forms of the hydroxide are formed. Whether or not the iron system would flocculate the low molecular weight residues without concomitant flocculation of polymers and/or suspended solids was not determined.

The types of carboxylic acids which might be derived from wood and form complexes with iron include the Krebs cycle acids (citric, adipic, oxalacetic, etc.), oxalic, aldonic, and uronic acids, and aromatic acids. Since little ultraviolet absorption was found associated with the low molecular

weight gel chromatography fractions, aromatic residues have not been considered in this phase of the study.

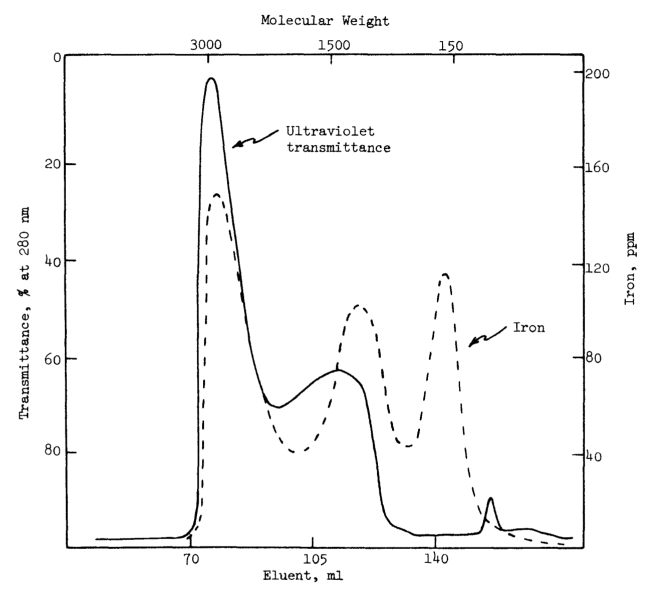


Figure 21. Transmittance and molecular weight distribution of a gel-chromatographed solution of sodium gluconate and ferrous ammonium sulfate, pH 7.0.

The sewer to the Combined Locks clarifier has five principal tributaries two of which gave aldose test responses strong enough for analysis by gel chromatography after the effluent was concentrated fivefold. The column elution curves are shown for the digester room effluent and the paper mill effluent (Figure 22). The digester room effluent showed predominantly high molecular weight residues and the paper mill effluent, low molecular weight residues.

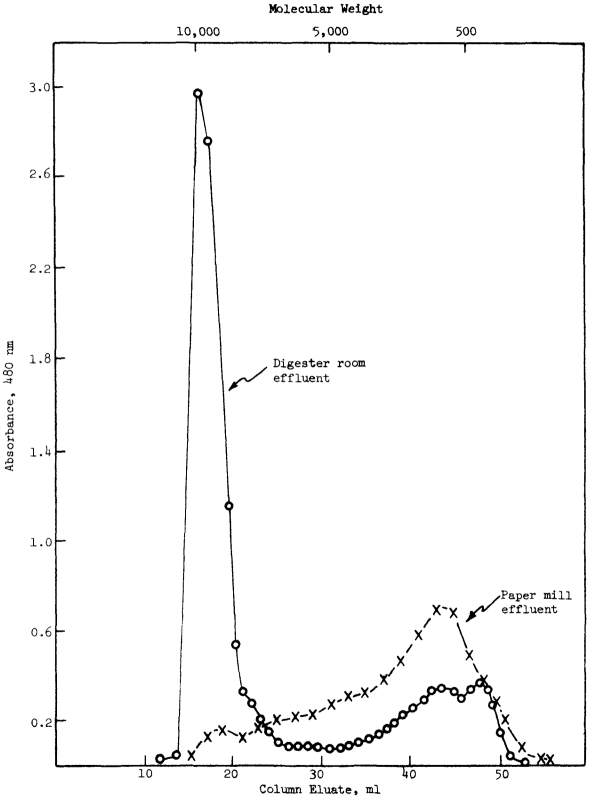


Figure 22. Phenol sulfuric acid test for aldoses and molecular weight distribution of gel-chromatographed chemimechanical pulp mill effluents.

Since this particular mill uses starch as one of its additives, one of the aldoses could be glucose and/or related starch degradation products.

STUDIES ON INFLUENT AND EFFLUENT SAMPLES FROM THE FIELD TRIALS

Clarifier influent and effluent samples from the two trials, at Mosinee and Combined Locks mills, were sent to the Analytical Section of the Institute for carbohydrate and weak acid determinations. Data are given in Table 40 for the results of these analyses on samples from high, intermediate and low soluble BOD₅ removal levels during clarification.

The carbohydrates were determined by gas chromatography after acid hydrolysis and derivative formation $(\underline{15})$. The weak acids were determined by passing the sample through a cation-exchange resin column and conductimetrically titrating it with 0.1N sodium hydroxide.

No consistent correlation was apparent between the soluble BOD_5 and the carbohydrate removals, indicating that in the high soluble BOD_5 removals some precursor(s), in addition to other than the oligo- and polysaccharides, were being removed by flocculation and clarification.

There was slight correlation between the soluble BOD₅ and weak acid removal values for these samples, particularly those from the Mosinee study.

STUDIES WITH MODEL COMPOUNDS

On a modest scale we screened model organic compounds for removal from solution by flocculation in order to learn more about the chemical nature of susceptible compounds.

The compounds were selected on the following basis:

- 1. Those compounds having at least some structural relation to substances found in mill effluents, either as atypical compounds or "families of compounds" (16-18);
- 2. Those compounds with some BOD, without marked "toxic" effects; and
- 3. Compounds having sufficient solubility so that gas chromatography, total organic carbon or some other convenient analytical procedure could be used to monitor the results of jar tests.
- 4. Removals of TOC from pure solutions were early recognized to be critically affected by pH. This variable was evaluated carefully and the interesting and probably important results provided in the following figures are discussed with the realization that the highest levels of TOC removal were often noted at pH levels outside the ranges normal for mill wastes or discharged to treatment systems.

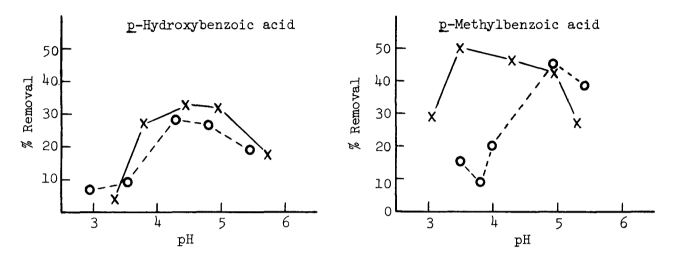
The jar tests were made with added fiber in the form of unbleached kraft pulp which had been disintegrated in an Oster blender for 5 minutes to provide a surface area for adsorptive bridging. Fibers had increased the removal of BOD₅ in some of our earlier work (Section V). This effect was confirmed with two runs with hydroxybenzoic acid and p-methylbenzoic acid with Fe³⁺

Table \downarrow 0. Removal of Carbohydrates and weak acids from mosinee and combined locks waste streams by Pilot-scale clarification^a

									24	Removal, %	
Total BODs, mg/l Influent Efflue	Ds, mg/l Effluent	Soluble BODs, mg/l Influent Effluent	DDs, mg/l Effluent	Carbohydrates, mg/l Influent Effluent	Effluent	Weak acids, meg/g ^b Influent Effluent	Ffluent	T	ပ္ကြက္က	Carbo- hydrates	Weak acids
Mosinee Samples	amples										
158	83	117	78	8.0	7.1	7.4	0.12	84	33	718	76
160	90	136	90	0.6	2. 6	2.1	0.16	†† †	34	143	95
148	122	108	92	5.7	8.7	9.4	3.0	18	15	-53	35
147	120	110	98	5.5	4.9	2.9	1.4	18	#	-16	52
150	136	112	108	15.8	4.8	1.8	1.8	9	4	70	0
1 6	92	42	16	4.8	4.6	5.9	1.2	a	†	8	59
Combined	Combined Locks Samples	Ø									
226	114	107	82	10.5	18.4	3.1	1.7	50	23	-75	45
180	48	104	78	22.9	31.0	1.5	1.2	53	25	-35	20
386	239	234	216	8.94	62.2	3.2	2.3	38	ω	-33	28
267	240	211	202	19.7	24.8	3.7	3.3	10	. †	-26	11

 $^{\rm a}_{\rm Samples}$ were from high, medium, and low BODs removal levels during clarification. $^{\rm b}_{\rm Milliequivalents/gram}$ of solids. $^{\rm c}_{\rm T}$, total; S, soluble.

(as FeCl₃) at various pH's, with and without the addition of fibers (Figure 23). TOC analyses were used to monitor changes in chemical concentrations of the model compounds before and after clarification.



- X With 350 mg/l kraft fiber added
- Without added fiber

Solutions were made from an amount of the model compound calculated to give a TOC level of 90-95 mg/l in the final volume to be used in the study. This was first dissolved in approximately 4 liters of distilled water; the correct amount of blended fiber was added (350 mg/l final volume), and the volume was brought to either seven or nineteen liters (final volume), depending upon whether one or two sets of jars were to be used for the evaluation.

One liter of the mixture containing the model compound and fibers was added to each of six jars. Five of the jars were treated with flocculating agent (various concentrations) and the sixth was used as an untreated "control." All jars were settled, sampled and filtered (0.45 μm). TOC was determined on the filtrates from the "control" and the five treated samples. All "removal" values are based on a comparison with the TOC concentration of the "control" after filtration. In this way the adsorption of the model compound on the fibers without flocculant addition could be monitored. In all tests the adsorption onto fibers of the controls was less than 8 percent (based on weight calculation), except for the arrowroot starch and dextrose which were found to be 65 and 24 percent, respectively.

Ferric chloride and alum were the primary flocculants used. Alum (Table 41) did not remove significant amounts of model compounds except polygalacturonic acid (87 percent removed at 200 mg/l alum concentration). The highest removal of model compounds obtained with lime up to 500 mg/l was 2 percent.

TABLE 41. TOC REMOVAL FROM SOLUTIONS OF MODEL COMPOUNDS TREATED IN JAR TESTS WITH ALUMINUM SULFATE

Compounds	Removal of TOC, %
3,4-Dihydroxycinnamic acid	0.0 ^e
Hydroxymalonic acid (tartronic acid)	1.0 ^e
Methylmalonic acid	1.0 ^e
D-Galacturonic acid	1.1 ^e
p-Hydroxybenzoic acid	2.2 ^e
Propanedioic acid (malonic acid)	2.6 ^e
p-Methylbenzoic acid	3.2 ^e
Dextrose	4.0 ^e
Isopentyl alcohol (isoamyl alcohol)	4.7 ^b
1,4-Butanedioic acid (succinic acid)	7.4 ^e
4-Hydroxy-3-methoxycinnamic acid (ferrulic acid)	8.3 ^e
3-Methoxy-4-hydroxy benzaldehyde (vanillin)	9.1 ^e
trans-Butenedioic acid (fumaric acid)	10.1 ^e
Arrowroot starch	11.4 ^d
p-Hydroxycinnamic acid	13.3 ^c
Benzylmalonic acid	34.4 ^e
Polygalacturonic acid	87.0 ^d

Only the best removals are listed in this table.

b Alum = 50 mg/l.

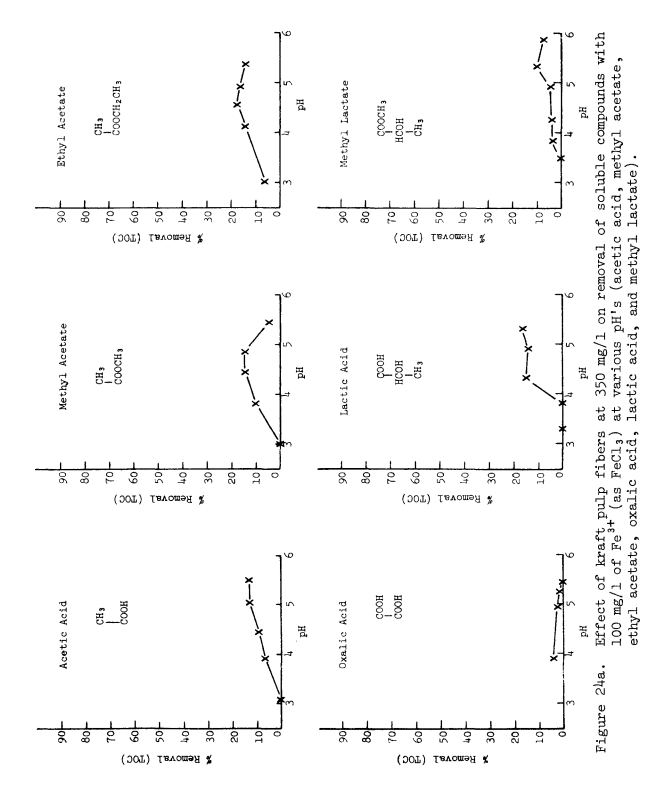
c Alum = 100 mg/l.

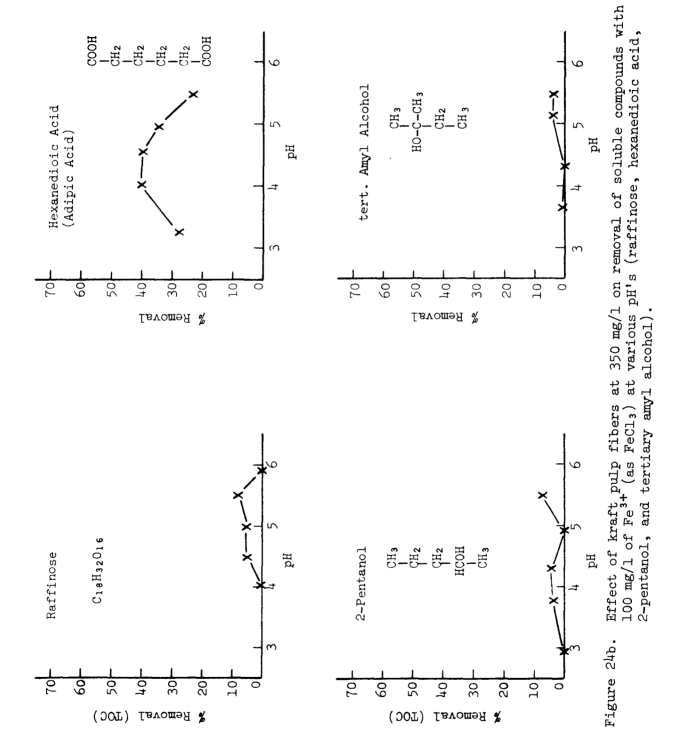
d Alum = 000 ^aAll compounds were treated with all levels of alum.

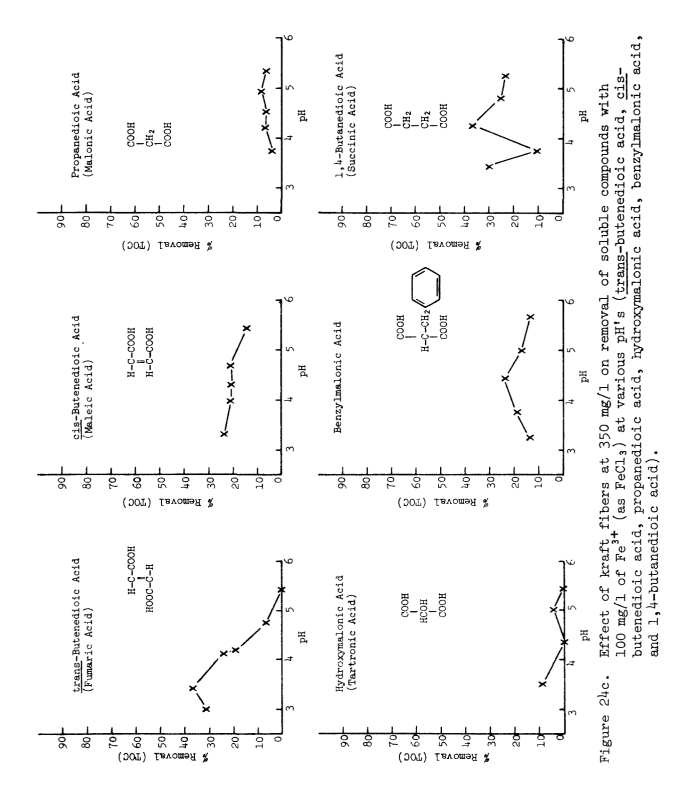
Data for the TOC removed when other compounds were similarly tested are given in Figure 24a-e. These data are very complex, with low molecular weight short-chained compounds (acetic acid, ethyl acetate) being removed at a 20 percent level, 2-pentanol and tertiary amyl alcohol showing less than 5 percent removal and hexanedioic acid, 40 percent removal. Several aromatic carboxylic acids had high removals, while galactouronic acid reductions were less than 5 percent (Figure 24e).

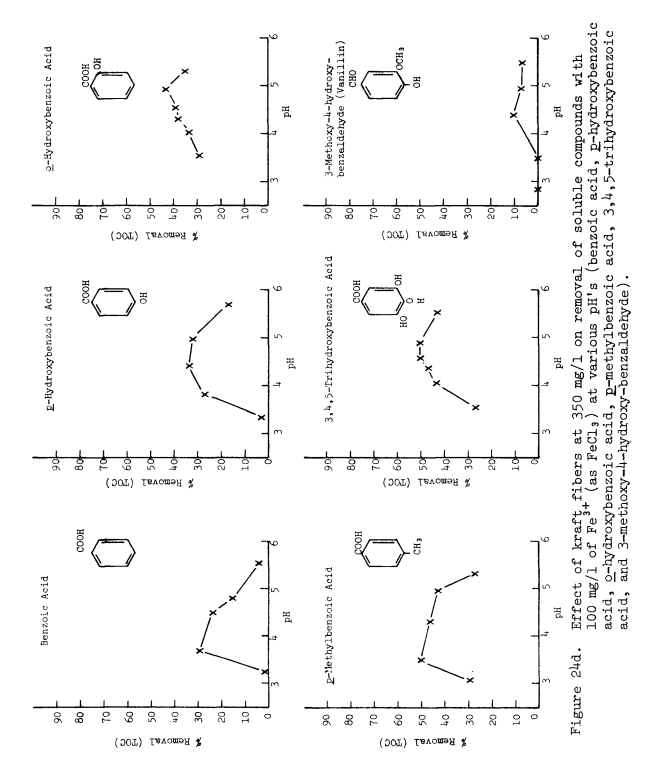
Alum = 200 mg/l.

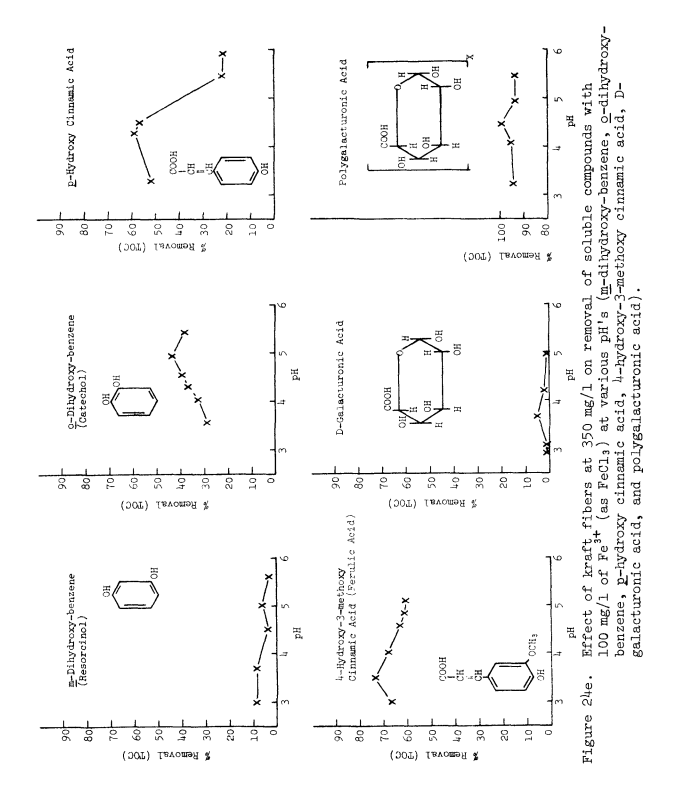
eAlum = 300 mg/l.











Some of the greater reductions in TOC could be traced to the formation of insoluble salts with the ferric ion, such as for the 3,4,5-trihydroxybenzoic acid (Figure 24d). This would not account, however, for the removal of p-hydroxybenzoic acid from solution since the ferric salt is soluble. Nor would the esters and alcohols, which do not form salts, be removed by this reaction.

The water solubility of the compounds, however, roughly correlates with their removal from solution with Fe³⁺ (Table 42). The compounds group into two classes, those with greater than 10 percent solubility in water and those with less than or equal to 1.5 percent solubility; the former group tends to have removals of 15 percent or less and those in the second class tend to have removals of 25 percent or greater (Figure 25).

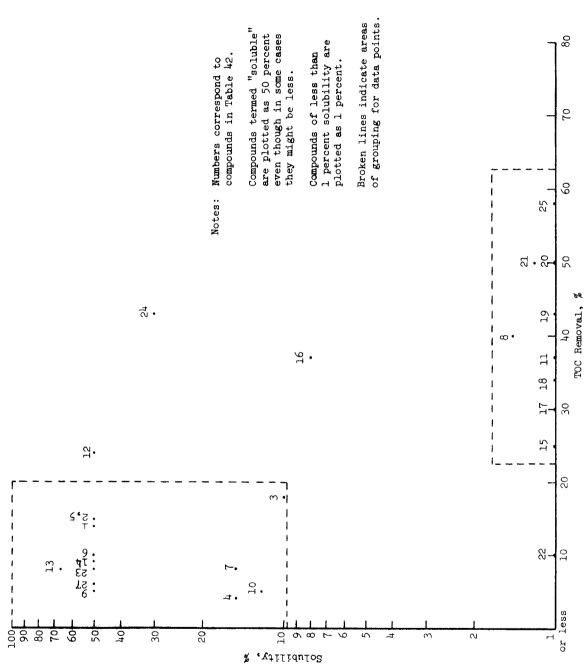
We must also consider not only reactions such as the cross-linking of starch phosphates with various cations and the increased complexing tendency observed for K⁺, Ca⁺⁺, Zn⁺⁺, Cu⁺⁺ and Fe⁺⁺⁺ that Wettstein, Neukom and Deuel $(\underline{19})$ studied extensively but also the oxalate or salicylate complexes with aluminum or iron as discussed by Stumm and Morgan $(\underline{14})$. Stumm and Morgan studied the formation of ferric-hydroxo-complexes and found that they tended to polymerize with other charged molecules at rather exact pH's. Such interactions might be occurring in our jar tests.

Additional work would be required to pinpoint the effects on soluble BOD_5 removal by the organic compound chain lengths or branching, the presence of phenolic or carboxylic groups and the solubility of the individual compounds, all of which appear to have some important relationship to the overall soluble BOD_5 removal mechanism.

TABLE 42. SOLUBILITY IN WATER OF VARIOUS MODEL COMPOUNDS AND THEIR REMOVAL IN JAR TESTS WITH Fe³⁺ AND PULP FIBERS^a

Reference No.	Compound ^a	Solubility, b %	Removal, a
1	Acetic acid	Miscible	14
2	Methyl acetate	Soluble	15
3	Ethyl acetate	10	18
4	Oxalic acid	15	14
5	Lactic acid	Miscible	15
6	Methyl lactate	Decomp. in H ₂ O	10
7	Raffinose	15	8
8	Adipic acid	1.44	40
9	2-Pentanol	Soluble	5
10	Tertiary amyl alcohol	12	5
11	Fumaric acid	0.63	37
12	Maleic acid	Freely soluble	24
13	Malonic acid	66	8
14	Tartronic acid	Very soluble	9
15	Benzylmalonic acid	Not given; probably	
		less than 1%	25
16	Succinic acid	8	37
17	Benzoic acid	0.3	30
18	<u>p</u> -Hydroxybenzoic acid	0.8	34
19	<u>o-</u> Hydroxybenzoic acid	0.2	43
20	<u>p-</u> Methylbenzoic acid	Slightly soluble	50
21	3,4,5-Trihydroxybenzoic acid	1.2	50
22	Vanillin	1	10
23	Resorcinol	50	, 8
24	Catechol	30	43
25	p-Hydroxycinnamic acid	Not given; probably less than 1%	58
26	Ferulic acid	?	73
27	D-galacturonic acid	Soluble	6
28	Polygalacturonic acid	?	99

Model compounds and % removal shown in Fig. 23-24e. bSolubility in water, from Merck Index, 7th edition.



The model compounds shown in Figures 23-24e cluster in two groups on the basis of solubility in water and removability with ${\rm Fe}^{3+}$. Figure 25.

SECTION IX

AREAS FOR COST REDUCTIONS

This research study on the mechanisms for the BOD_5 removal in clarifiers was not intended to cover comparative process economics. However, evaluation of the data obtained in the laboratory and practical observations in the field studies indicate that improvements in the clarification process will reduce costs. Studies on the flows from individual process sewers demonstrated that treatment which removes soluble BOD_5 prior to the clarifier is likely also to reduce the overall cost for the total system of treatment.

SEPARATION OF FLOWS ADVERSELY AFFECTING CLARIFICATION EFFICIENCY

There are several ways to reduce costs, and careful control of the quality of the in-flow to the clarifier is one of them. Proper clarifier performance (reduction of suspended solids, COD, BOD5, etc.) is dependent upon sedimentation and flocculation properties of the influent flow. The input of lignin and other pulping and bleaching constituents having colloidal and dispersant properties is of particular concern. The reduction or elimination of flows containing such materials, and particularly the containment and separate treatment of large spills of them, should substantially improve the clarification efficiency and should have high priority in programs for cost reduction in waste treatment. Revised handling should produce the following substantial economies:

- 1. The capital investment for a clarification system will be less when the clarifier size is reduced;
- 2. Reagent and other operating charges will be less as clarifier efficiency improves;
- 3. Clarifier performance in all categories, including the amount of BOD₅ removal, should be greatly improved;
- 4. Although no measurements were undertaken in this study, sludge dewatering, sludge volume and sludge disposal problems were apparent and should be subject for further development;
- 5. Needs and costs for subsequent (secondary biological) treatment may be significantly reduced in terms of capital cost and operating charges, and the need for secondary clarification may also be reduced.

OPTIMIZING THE TYPE AND AMOUNT OF CHEMICAL ADDITIVE

This project also showed the importance of optimizing the amounts and types of flocculating agents for various clarifier inflows. Different chemical additives are needed for different mills. Process fluctuations and changes affecting clarifier performance were apparent within the short term of each mill field trial.

Operating experience may make possible the development of routines for addition of chemicals when the effluents change. The rising trend toward development and use of computerized systems to control the manufacture of pulp and paper may logically extend to automated changes during effluent treatment.

Of the two mill clarifiers used in the field studies, one was utilizing chemical additives routinely and the other clarifier was well equipped to use flocculating chemicals. Capital costs for additional chemical feeding and mixing equipment would not be a major expense. Operating charges for the chemical additives that we used was about 3ϕ to 10ϕ per 1000 gallons of clarifier inflow (Table 43). Since these chemicals might logically replace less efficient additives already being used, cost could be considerably less in actual practice. The efficient use of chemicals would, moreover, be expected to give a return in reduced overall waste treatment costs. The actual cost of chemical additives to remove soluble BOD₅ cannot be adequately assessed short of detailed engineering cost studies to fit the situations in each mill.

TABLE 43. CHEMICAL COSTS FOR CLARIFICATION

Type of flocculant	Required, mg/l	Cost, ^a ¢/1000 gallons
Fe ^{3+b}	25	2.72
	50	5.44
	100	10.88
Alum ^c	100	3.13
	200	6.26
	300	9.39

Addition of 0.75 mg/l Hercofloc 812.3 would add 0.5¢.

^bAs ferric chloride (FeCl₃·6H₂O) at \$90/ton.

CAluminum sulfate •16 H₂O at \$75/ton.

SECTION X

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SECTION XI

GLOSSARY

- Alum Aluminum sulfate.
- BOD Biochemical oxygen demand is based on the oxygen requirement of living organisms while utilizing components of the waste stream for growth and/or reproduction.
- BOD₅ The biochemical oxygen demand in a 5-day test period at 20°C.
- <u>CM</u> Chemimechanical pulping; this is based on a short chemical cook followed by mechanical refining to separate the fibers in the softened chips.
- <u>COD</u> Chemical oxygen demand is the measurement of the oxygen equivalent of that portion of the organic matter in a sample that is susceptible to oxidation by strong chemical oxidants (e.g., chromic acid).
- <u>Colloid</u> A phase dispersion to such a degree that the surface forces become an important factor in determining its properties. General particles of colloid dimension are 0.001 to 1 micron.
- Gel chromatography A column chromatographic technique which fractionates molecules on the basis of molecular weight.
- Hemicellulose That fraction of plant stems which is made up of carbohydrate
 polymers other than cellulose.
- Molecular exclusion The mechanism of fractionation of molecules during gel chromatography. Larger molecules are restricted from diffusing into the gel matrix, thus taking less time to travel through the gel than smaller molecules.
- <u>Permeate</u> That portion of a solution passing through a membrane during reverse osmosis or ultrafiltration.
- Reverse osmosis (RO) Osmosis in reverse flow through a semipermeable membrane when external pressure in excess of the osmotic pressure is applied.
- Soluble As used in this report is that portion of the solution (solvent and solute) passing through a 0.45 μm filter under 100 psig nitrogen pressure.
- Zeta potential That portion of the total potential drop between the surface of the solid and the suspending liquid that is contributed by the charge potential between the liquid adhering to the wall of the particle and the movable liquid.

SECTION XII

APPENDICES

APPENDIX A: COMPANIES AND AGENCIES SUPPORTING PHASE I STUDIES

Bergstrom Paper Company

Hammermill Paper Company

Hoerner Waldorf Corporation

Kimberly-Clark Corporation

The Mead Corporation

NCR-Appleton Papers Division

Nekoosa Edwards Paper Company, Inc.

Potlatch Corporation

The Proctor & Gamble Company

Scott Paper Company

Wausau Paper Mills Company

Department of Natural Resources, State of Wisconsin U.S. Environmental Protection Agency

TABLE B-1. CLARIFIER STUDIES WITH HERCOFLOC 812.3 (0.75 mg/1) AT MOSINEE PAPER CORPORATION

					Influent (A)	(A)			ഥ	Effluent (B)	(B)			% Reduction	tion	
Date	Day	Composite	Hd	Sus. Sol., mg/1	, BOD5, Total	, mg/l Soluble	Color Units	Hd	Sus. Sol., mg/1	12	BOD5, mg/l tal Soluble	Color Units	Sus. Sol,	Total	BOD5 Soluble	Color
	ue to	Due to Mechanical Difficulties - No "Se	ifficul	ties - No	ts of	Samples" we	were taken	from 1/15	15 - 2/5/75	(lst to	22nd day)					
2/6	23	1 2 M M11 ^b	7.70	167 202 196 218	96 127 116 90	72 120 96 	45 93 	7.59 7.90 7.61	45 50 41 56	85 120 94 51	88 108 87	52 97 52	73.0 75.2 79.1 74.3	11.4 5.5 19.0 43.3	-22.2 10.0 9.4	-13.5 -4.1 46.4
7/2	24	1 2 M Mill ^b	7.90 7.81 7.52	273 211 266 260	111 137 132 84	110 131 123	93 103 82 	7.91 7.82 7.61	74 45 44 56	121 113 113 63	106 92 99	74 74 63	72.9 78.7 83.4 78.5	-9.0 17.5 14.4 25.0	3.6 29.8 19.5	20.4 28.2 23.2
2/11	28	1 2 M M111 ^b	7.60 7.50 9.20	358 359 257 249	372 177 262 249	179 142 254 	374 86 1580	9.73 8.40 9.09	101 153 53 71	365 149 222 195	246 150 217	5040 118 1420	71.8 57.4 79.4 71.5	-1.9 15.8 15.3 21.7	-37.4 -5.6 14.6	-124.7 -37.2 10.1
2/12	29	1 2 M M111 ^b	7.39 7.70 8.40	416 344 502 338	152 188 174 141	122 142 166 	50 144 38	7.40 6.88 7.59	106 97 88 66	134 143 141 114	127 124 144 	45 56 81	74.5 71.8 82.5 80.5	11.8 23.9 19.0 19.1	-4.1 12.7 13.2	10.0
2/13	30	1 2 M Mill ^b	8.53 8.69 8.21	364 256 290 160	138 204 152 111	120 162 129 	56 284 104 	8.29 8.38 7.89	114 131 85 73	124 162 130 108	126 142 120	63 168 115	68.7 48.8 70.5 54.4	10.1 20.6 14.5 2.7	-5.0 12.3 7.0	-12.5 40.8 10.6
2/14	31	1 2 M Mill b	8.40 8.21 8.12	228 266 289 210	170 150 128 99	114 127 97 	63 48 86 	8.12 8.19 7.70	73 49 37 57	111 120 124 75	114 125 110 	49 38 59	68.0 81.6 87.2 72.8	34.7 20.0 2.9 24.2	-13.4	22.2 15.6 31.4

**Composites
A1 - from 7:30 a.m. to 7:30 p.m.
A2 - from 7:30 p.m. to 7:30 a.m.
B1 and B2 - Two hours after A's to allow for holding time
M - 24-hour mill composite (7:30 AM) analyzed in Appleton labs.

**Same as (M) analyzed at mill.

TABLE B-2. CLARIFIER STUDY WITH ALUM (200 mg/l) PLUS HERCOFLOC 812.3 (0.75 mg/l) AT MOSINEE PAPER CORPORATION

				Inf	13	(A)			B	Effluent (B)	8)			% Reduction	tion	i
Date	Day	Composite ^a	Hd	Sus. Sol., mg/l	BOD5, mg/1 Total Solu	mg/1 Soluble	Color Units	Нd	Sus. Sol., mg/1	BOD5, Total	mg/l Soluble	Color Units	Sus. Sol.	BOD5 Total Sc	D5 Soluble	Color
2/18	35	1 2 M M11 ^b	9.40 8.78 9.29	188 54 342 464	112 130 128 81	108 106 127	108 74 652	8.01 5.80 8.58	10 8 61 64	78 77 74 66	80 85 76	4 0 97	94.7 85.2 82.2 86.2	30.4 40.8 42.2 18.5	25.9 19.8 40.2	96.3 100.0 85.1
2/19	36	1 2 M M111 ^b	8.51 9.11 8.80	213 222 218 234	146 190 150 99	128 174 112	203 888 356	7.32	0.5 63 74	104 108 136 69	98 107 108	19 41 188	98.1 99.9 71.1 68.4	28.8 43.2 9.3 30.3	23.4 38.5 3.6	90.6 95.5 47.2
2/20	37	1 2 M M11 ^b	7.30 8.11 8.41	119 136 153 216	103 139 116 159	73 117 99	80 310 170	8.55 6.80 7.78	15 18 62	92 90 102 135	94 88 78	22 15 125	87.4 86.8 73.8 71.3	10.7 35.2 12.1 15.1	24.8 24.8 21.2	72.5 95.2 26.5
2/21	38	1 2 M Mill ^b	7.83 8.08 7.59	278 289 206 204	116 140 112 129	112 127 88	71 133 56	8.19 7.62 7.39	44 70 55 58	104 126 104 108	99 106 87	52 74 74 	84.2 75.8 73.3 71.6	10.3 10.0 7.1 16.3	11.6	26.8 44.4 -32.1
2/26	43	1 2 M Mill ^b	7.41 7.58 7.54	221 176 215 283	114 113 140 114	112 94 96 	41 23 31	8.84 8.71 7.40	60 7 60 79	100 113 104 105	98 84 1-	4 0 15	72.8 96.0 72.1 82.2	12.3 0 25.7 7.9	16.1 4.2 8.3	90.2 100.0 51.6
2/27	77	1 2 M M11b	7.55 7.78 7.65	155 153 180 248	108 100 110 96	108 104 104	120 122 124 124	6.90 6.68 7.25	24 18 62 57	96 87 86 75	9 8 9 1 9 3 5 1	22 12 78	84.5 88.2 65.6 77.0	11.1 13.0 21.8 21.9	14.8 18.3 10.6	81.7 90.2 37.8
2/28	45	1 2 M Mill	7.53 7.70 7.32	82 130 201 121	96 94 86 108	95 89	62 68 68	6.55 6.20 7.25	9 15 71 30	84 83 78 36	83 80 71	22 12 49	89.0 88.5 64.7 75.2	12.5 11.7 9.3 66.7	12.6	64.5 85.0 27.9
ď	1 4 5 5															

**Composites
Al - from 7:30 a.m. to 7:30 p.m.
A2 - from 7:30 p.m. to 7:30 a.m.
Bl and B2 - Two hours after A's to allow for system holding time
M - 24-hour mill composite (7:30 AM) analyzed in Appleton labs.
**Mill - same as (M) analyzed by mill.

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TABLE B-3. CLARIFIER STUDY WITH FERRIC CHLORIDE (100 mg/1 Fe3+) PLUS HERCOFLOC 812.3 (0.75 mg/1) AT MOSINEE PAPER CORPORATION

				q_	Influent (A)				Rf	Effluent (B)	£			% Reduction	ction	
Date	Day	Composite	ЬĦ	Sus. Sol., mg/l	BOD5, otal	mg/l Soluble	Color Units	Нd	Sus. Sol., mg/1	1.4	mg/l Soluble	Color Units	Sus. Sol.	BC Total	BOD5 Soluble	Color
3/4	67	1 2 M M113	7.85 8.75 8.00	189 136 204 230	190 244 226 135	93 138 102 	78 130 162 	4.10 3.93 7.95	18 30 74 66	70 124 110 72	72 106 84	45 30 140 	90.5 77.9 63.7 71.3	63.2 49.2 51.3 46.7	22.5 23.2 17.6	42.3 76.9 13.6
3/5	50	1 2 M M111 ^b	7.73 8.51 7.62	126 199 207 190	128 158 148 135	88 117 108	133 204 204 	3.74 2.69 7.71	28 83 37 42	86 83 122 105	78 78 92	34 27 156	77.8 58.3 82.1 77.9	32.8 47.5 17.6 22.2	11.4 33.3 14.8	74.4 86.8 23.5
3/6	51	1 2 M M11 ^b	8.40 7.61 7.80	240 169 276 264	140 142 122 107	95 88 91	168 74 98	3.47 3.18 7.28	22 6 63 49	86 88 106 77	75 76 82 	27 12 74	90.8 96.4 76.9 81.4	38.6 38.0 13.1 28.0	21.0 13.6 9.9	83.9 83.8 24.5
3/7	52	1 2 M Mill ^b	7.11 7.31 7.22	198 213 204 224	88 94 91 105	86 76 74	82 56 100	3.89 3.89 7.18	4 75 65	80 92 96 81	78 79 70	27 19 60	98.0 99.1 63.2 71.0	9.1 2.1 -7.0 22.8	9.3 -3.9	67.1 66.1 40.0
3/11	95	1 2 M b Mill ^b	8.89 7.81 7.70	182 209 197 264	102 114 102 111	82 98 81	170 60 1111	3.89 4.18 7.41 	8 5 51	73 78 109 93	99 77 87 	15 19 118	95.6 97.6 67.0 80.7	28.4 31.6 -6.8 16.2	19.5 21.4 -7.4	91.2 68.3 -6.0
3/12	57	1 2 M M11b	7.69	228 22 258 236	170 109 130 132	144 102 100 	126 38 71 	3.50 4.39 7.31	9 4 45 39	120 70 94 81	79 71 94	30 27 38	96.0 81.8 82.6 83.5	29.4 35.8 27.7 38.6	45.1 30.4 6.0	76.2 28.9 46.5
3/13	58	1 2 M Mill	8.26 7.99 7.82	220 199 206 294	160 161 140 141	136 127 114	192 174 222	4.71 4.11 7.51	12 13 81 57	90 100 121 126	90 99 112	52 52 122	94.5 94.0 60.7 80.1	43.8 37.9 13.6 10.6	33.8 22.0 1.8	72.9 70.1 45.0
3/14	59	1 2 M Mill ^b	7.10 7.29 7.41	192 128 245 176	122 128 142 144	101 102 110	115 159 126	4.18 3.51 7.19	15 4 59 59	92 86 129 108	83 81 110	38 34 108 	92.2 96.9 75.9 66.5	24.6 32.8 9.2 25.0	17.8 20.6 0	67.0 78.6 14.3
8	:															

**Composites - all on 60-minute cycle
Al - from 7:30 a.m. to 7:30 p.m.
A2 - from 7:30 p.m. to 7:30 a.m.
Bl and B2 - Two hours after A's to allow for system holding time
M - 24-hour mill composite analyzed at Appleton labs.

**Mail - same as (M) analyzed by mill.

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CLARIFIER STUDY WITH FERRIC SULFATE AND HERCOFLOC AT MOSINEE PAPER CORPORATION	
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				In	fluent (A)			Ed	Effluent (B)				% Reduction	ion	
Date	Day	${\tt Composite}^{\tt B}$	Hd 1	Sus. Sol., mg/1	BOD5, Total	mg/1 Soluble	Color Units	Hd	Sus. Sol., mg/1	BOD5, Total	mg/1 Soluble	Color Units	Sus. Sol.	BOD5 Total Sc	Soluble	Color
D. F.	erric	Ferric Sulfate (50-100 mg/1	100 mg/1	[Fe ⁺³)												
3/18	63	1 2 M Mill	8.01 7.28 7.41	187 214 359 246	192 147 221 195	138 110 131	332 82 126	6.16 6.32 7.30	126 105 57 51	124 120 144 93	101 98 132	38 38 211	32.6 51.4 84.1 75.2	35.4 18.4 34.8 52.3	26.8 10.9 -0.8	88.6 53.6 -67.5
3/19	99	1 2 M Mill	7.62	286 202 178 226	158 136 146 122	130 116 113	118 111 108 	5.30 4.56 7.01	22 18 73 60	100 93 121 110	95 88 106 	38 45 163	92.3 91.1 59.0 73.4	36.7 31.6 17.1 9.8	26.9 24.1 6.2	67.8 59.4 -50.9
3/20	65	1 2 M Mill ^b	8.09 7.22 7.07	191 168 149 175	180 173 186 140	148 124 144	273 145 163	4.67 3.61 7.11	55 10 74 59	122 86 140 86	107 83 122	45 52 160	71.2 94.0 50.3 66.3	32.2 50.3 24.7 38.6	27.7 33.1 15.3	83.5 64.1 1.8
3/21	99	1 2 M M b	7.40	200 178 224 218	165 139 170 120	135 104 116 	122 68 90	3.96 3.32 7.19	8 7 62 54	106 89 126 60(?)	98 86 106 	55 70 90	96.0 96.1 72.3 75.2	35.8 36.0 25.9 50.0(?)	27.4 17.3 8.6	54.9 -2.9 0
н М	ase Li	Base Line (Hercofloc 812.3		at 0.75 mg/1)	~											
3/25	70	1 2 M Mill ^b	11.01 7.91 8.95	341 190 241 278	119 134 132 93	91 104 100	52 31 56	11.02 7.81 8.26	68 23 61 63	102 112 88 63	90 112 86 	45 31 52	80.0 88.0 74.7 77.3	14.3 16.4 33.3 32.2	1.1 -7.7 14.0	13.5 0 7.1
3/26	71	1 2 M Mill ^b	7.99 8.06 7.81	239 182 262 300	170 138 	146 127 114 	41 34 18	7.80 7.78 7.51	36 24 46 50	170 112 125 47	114 112 112	41 34 38	84.9 86.8 84.0 83.3	0 18.8 48.9	21.9 11.8 1.8	000
3/27	72	1 2 M Mill ^b	8.32 8.13 8.19	137 235 245 192	172 146 192 108	130 130 122	103 192 150	8.41 8.14 7.84	51 39 57 51	182 125 132 57	128 124 104	100 296 103	62.8 83.4 76.7 73.4	-5.8 14.4 31.2 47.2	1.5 4.6 14.8	2.9 -54.2 31.3
3/28	73	1 2 M Mill ^b	8.28 8.51 9.21	177 189 253 260	184 150 158 117	127 134 120 	332 328 318 	8.01 7.99 8.32	31 48 62 43	166 125 120 120	139 124 110 	460 251 278 	82.5 74.6 75.5 83.5	9.8 16.7 24.0 -2.6	-9.4 7.5 8.3	-38.6 23.5 12.6
a Compo	osites	, - Al, A2, B	M. and !	Composites - Al, A2, Bl, and B2 on 30-mini	te cvele											

*Composites - Al, A2, Bl, and B2 on 30-minute cycle
Al - from 7:30 a.m. to 7:30 p.m.
A2 - from 7:30 p.m. to 7:30 a.m.
Bl and B2 - 90 minutes allowed for system holding time
M - 24-hour mill composite analyzed at Appleton labs.
Mill - same as (M) analyzed by mill.

TABLE B-5. CLARIFIER STUDY WITH SULFURIC ACID, SULFURIC ACID PLUS POLYMER, AND PERROUS SULFATE PAPER CORPORATION

	4			In:	nfluent (A)	.A.)			E	Effluent (B)	B)			% Reduction	tion	
Date	Day	Composite	ЬН	Sus. Sol., mg/l		mg/1 Soluble	Color Units	Hd	Sus. Sol., mg/1	BOD5, Total	mg/1 Soluble	Color Units	Sus. Sol.	Total	BOD5 Soluble	Color
F.	ulfuric	Sulfuric Acid (pH adjustment only)	justmen	t only)												
4/1	77	1 2 M b Mill	8.61 8.53 9.72	270 332 307 302	320 201 211	176 151 177	640 432 960 	6.51 6.19 9.51	193 113 50 62	248 143 140	150 112 125 	92 52 318	28.5 66.0 83.7 79.5	22.5 28.8 33.6	14.8 25.8 29.4	85.6 88.0 66.9
4/2	78	п	7.61	331	148	124	29	6.18	131	114	86	41	60.4	23.0	21.0	38.8
.5 SI	ulfuric	Sulfuric Acid + Polymer	mer													
	78	2 M Mill ^b	7.91	294 260 266	120 114 	102	74 59 	5.65	57 68 36	88 109 	76 92 	23 81 	80.6 73.8 86.5	26.7	25.5	68.9
н.	errous	Ferrous Sulfate (100 mg $\mathrm{Fe}^{+2}/1$) + Polymer	mg Fe+	$^{2/1}$) + Polyr	ner											
4/3	42	1 2 M M 11	7.70 8.72 8.19	271 177 227 224	150 144 140	126 114 120 	160 296 200 	5.19 7.61 7.30	183 169 83 52	106 148 104 	92 106 96	322 45 168 	32.5 4.5 63.4 76.8	29.3 -2.8 25.7	27.0 7.0 20.0	-101.2 84.8 16.0
4/4	80	1 2 M Mill ^b	8.61 7.62 7.40	222 184 275 356	172 138 216 	137 112 136	3600 200 118 	5.34 5.59 7.19 	117 164 57 45	144 90 143 	94 78 108 	136 207 185	47.3 10.9 79.3 87.4	16.3 34.8 33.8	31.4 30.4 19.1	96.2

**Composites - Al, A2, Bl, and B2 on 30-minute cycle
Al - from 7:30 a.m. to 7:30 p.m.
A2 - from 7:30 p.m. to 7:30 a.m.
B1 and B2 - 90 minutes allowed for system holding time
M - 24-hour mill composite analyzed at Appleton labs.
**Mill - Same as (M) analyzed by mill.

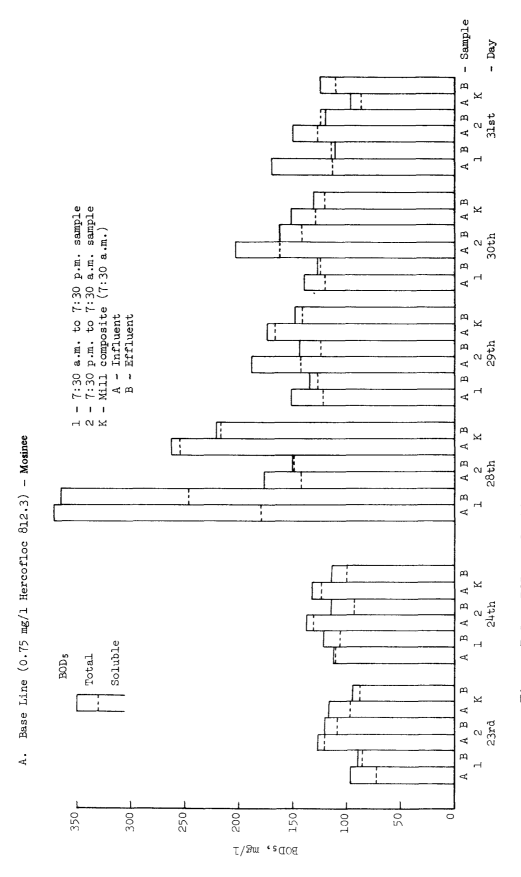
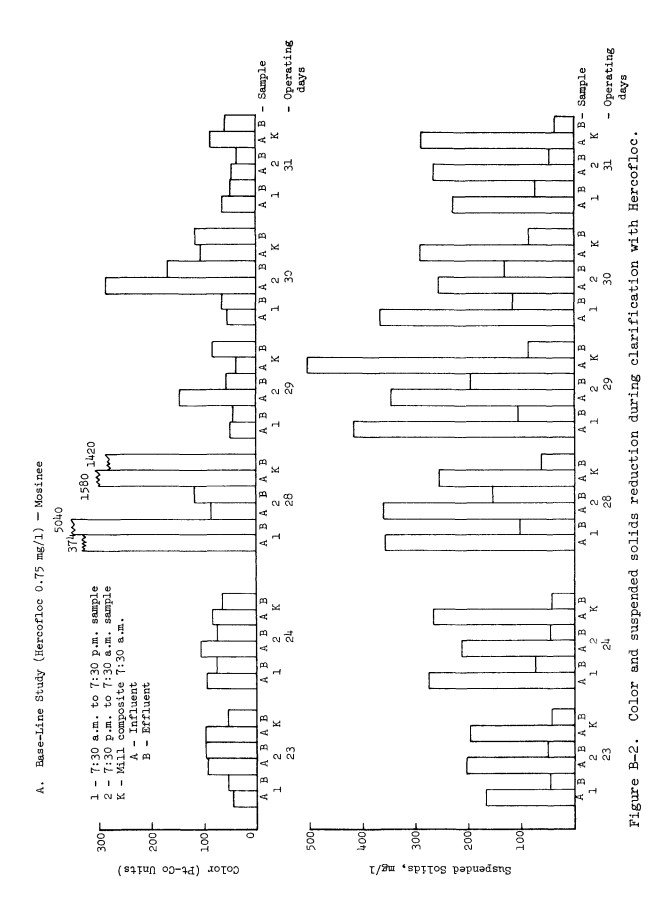


Figure B-1. BODs reduction during clarification with Hercofloc



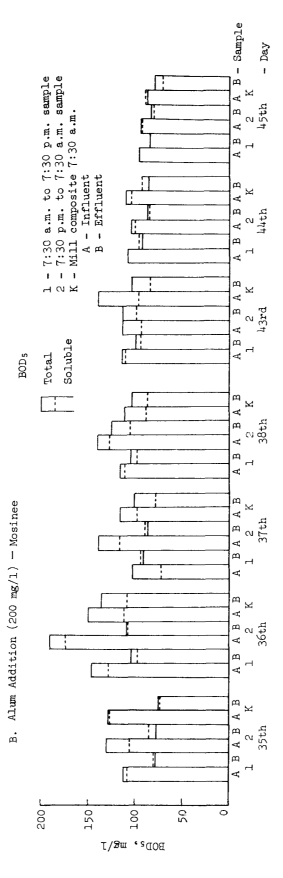
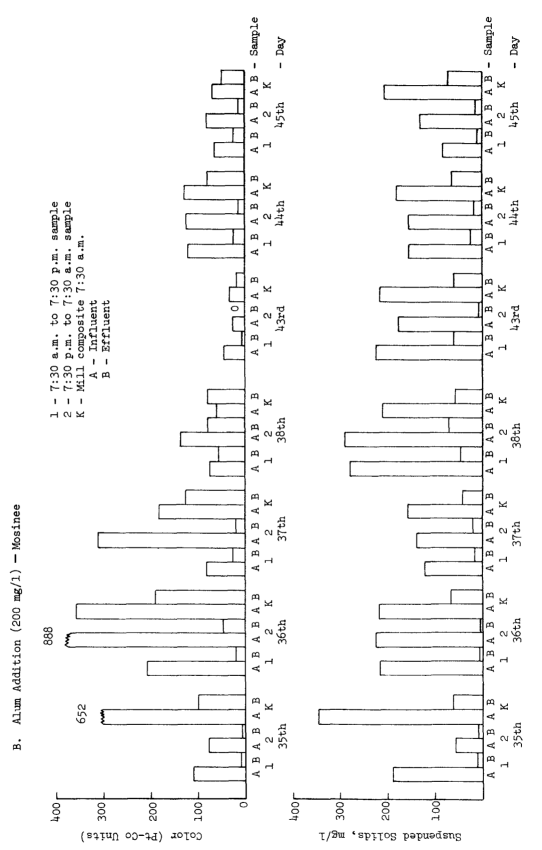


Figure B-3. BODs reduction during clarification with alum.



Color and suspended solids reduction during clarification with alum. Figure B-4.

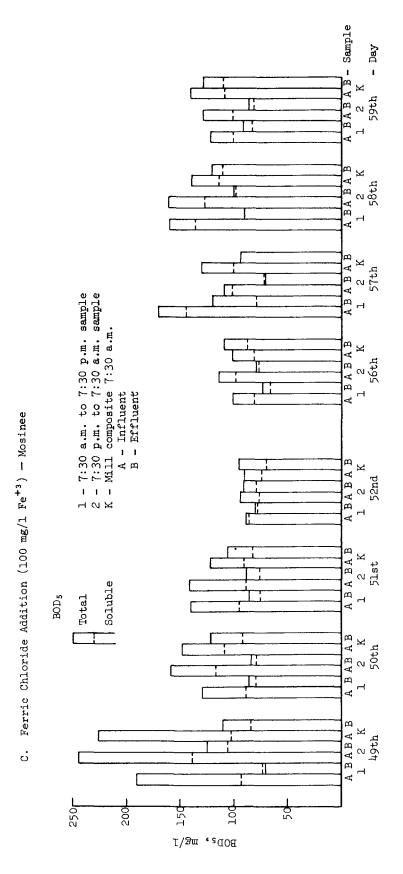
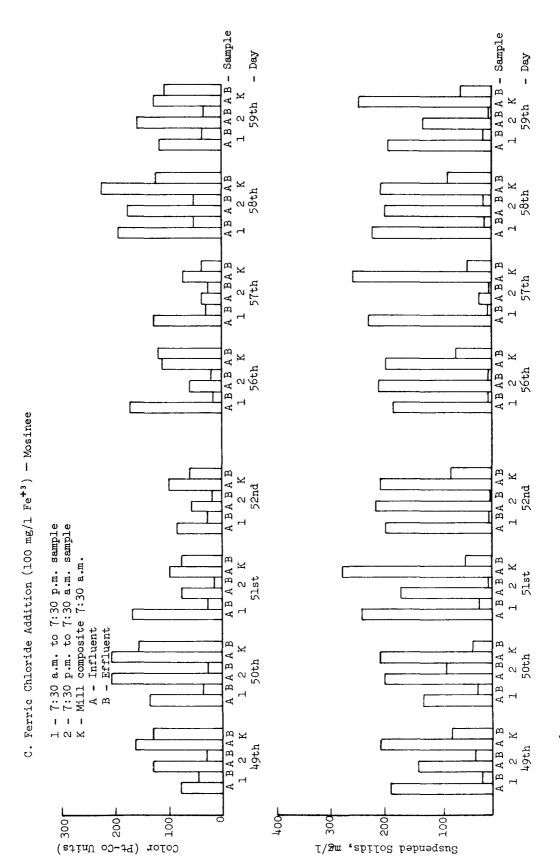
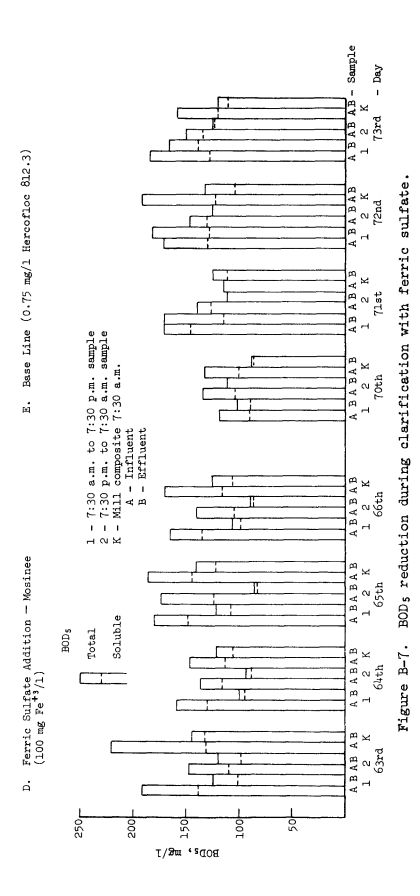
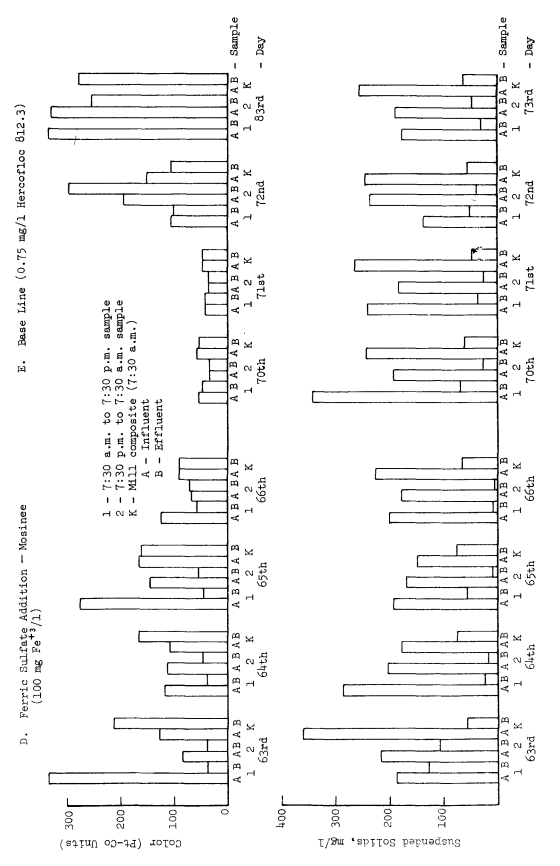


Figure B-5. BODs reduction during clarification with ferric chloride.

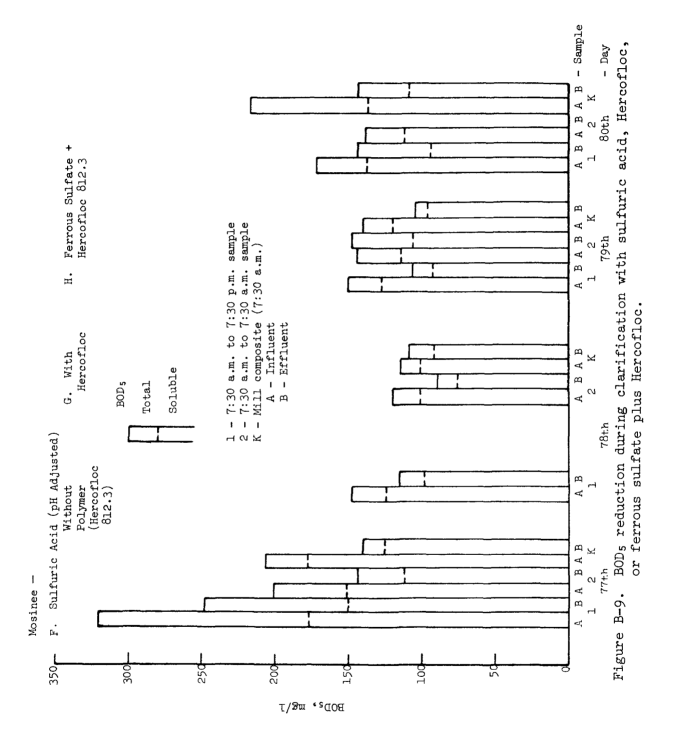


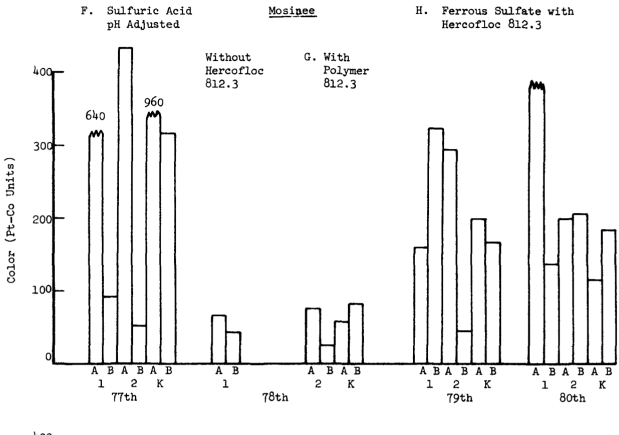
Color and suspended solids reduction during clarification with ferric chloride. Figure B-6.

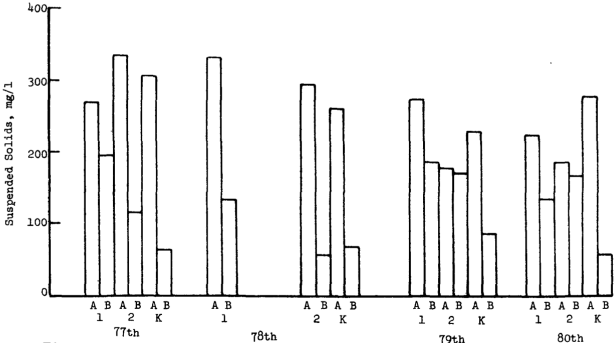




Color and suspended solids reduction during clarification with ferric sulfate. Figure B-8.







77th 78th 79th 80th
Figure B-10. Color and suspended solids reduction during clarification with sulfuric acid, Hercofloc, or ferrous sulfate plus Hercofloc.

TABLE C-1. CLARIFIER STUDY AT COMBINED LOCKS - BASE LINE (GRAVITY CLARIFICATION AFTER 0.010-INCH SIEVE) WITHOUT ADDITIVES^E,^b

				Influent (C)	t (C)						Efflu	Effluent (D)					%	% Reduction	uo	
Day	Composite	Hd	Sus. Sol. mg/l		BOD5, m Total So	mg/1 Soluble	Color Units	Sol. Sol. mg/l	Hd	Sus. Sol. mg/l		BODS, Total	mg/l Soluble	Color Units	Sol. Sol. mg/l	Sus. Sol.	01.	BOD5 Total Sc	Ds Soluble	Color
'n	2	8.30	805	25	594	196	162		7.71	76		256	201	192		88.0		12.9	-2.6	-18.5
10	40 E	7.65	578 749 664	4 E M	243 312 278	124 190 157	98 122 110	999	7.82 7.55 7.91	19 54 54		162 208 234	140 176 159	105 150 145	680 596	96.7 94.1 91.9		33.3 33.3 15.8	-12.9 7.4 0	-7.1 -22.9 -31.8
::	H ON E	7.58	758 814 786	2 8 21	267 322 294	202 233 218	125 215 170	796	7.62 7.71 7.69	66 37 48		240 282 236	211 238 209	217 202 255	850 740	91.3 95.4 93.9		10.1 12.4 19.7	-4.4 -2.4 4.1	-73.6 -40.5 -50.0
12	X 7 1	7.35	621 (3 547 (3 584 (3	(366) 17 (330) 20 (348) 15	176 204 190	127 142 134	100 112 106		7.50 7.67 7.38	19 (18 (29 ((16) (14) (22)	140 162 147	128 154 135	75 83 111		96.9 96.7 95.0	(95.6) (95.8) (93.7)	20.4 20.6 22.6	0 7.8- 0	25.0 26.0 -4.5
13	3 01	7.65	638 (4 643 (4 640 (4	427) 3 427) 3 427) 3	352 355 354	268 252 260	175 185 180	838	7.65 44 7.71 31 No Sample		(34) (27)	270 286	254 271	200	850	93.1 95.2	(92.0) (93.7)	23.3 19.4	5.2	-14.3 -9.8
14		7.55	584 (3	76 (292)	373	254	183		7.58	38	(25)	243	250	255		93.5	(93.7)	34.8	1.6	-39.3
17	H O E	7.40	541 (4 599 (4 570 (4	(403) 36 (426) 36 (426) 33	300 363 ·	216 232 <u>224</u>	195 187 191	796	7.40 7.50 7.32	72 (49 (55 ((44) (31) (32)	222 242 209	220 214 198	230 295 255	747 754	86.7 91.8 90.4	(89.1) (95.3) (92.5)	26.0 33.3 37.0	-1.8 7.8 11.6	-17.9
18	4 2 2	7.60	506 (3 622 (4 564 (4	(484) 23 (428) 37	236 377 306	148 286 217	126 218 172		7.68 7.70 7.55	31 55 51	(23) (41) (36)	164 277 201	147 241 190	155 244 292		93.9 91.2 91.0	(93.8) (91.5) (91.6)	30.5 26.5 34.3	0.6 15.7 12.4	-23.0 -11.9 -69.8
19	4 0 4	7.59	415 (2 435 (3 425 (3	(292) 21 (339) 36 (316) 25	217 380 298	138 319 228	204 322 263	966	7.58 7.59 7.60	36 ((30) (68) (42)	181 374 268	162 338 238	196 377 302	1040 798	91.3 82.5 88.0	(89.7) (79.9) (86.7)	16.6 1.6 10.1	-17.4 -5.6 -4.2	3.9 -17.1 -14.8

Ayalues in parentheses are volatile suspended solids. Underscored values are based on average of IPC influent samples.

TABLE C-2. CLARIFIER STUDY AT COMBINED LOCKS - FERRIC CHLORIDE (75 mg/l Fe³⁺) WITHOUT POLYMER

1	.	ł		_		_		_												_		
	Color		50.0		27.0	-11.3	14.2	20.6	1.0.1	29.2	-21.1		11.1	20.3	1	53.2	67.5	-39.8	44.1	55.9	-61.3	
uo	BODs Soluble		14.4 27.6	,	24.1 17.4	-2.6	27.6	20.0	y. y	31.0	9.1		31.2	22.6	10.0	28.6	32.4	22.8	23.4	35.0	6.7	
% Reduction	Total		37.5	0.07	48.3	3.8	49.7	42.1	72.7	35.0	19.5		47.0	21.0	7.67	45.2	45.6	36.2	9.67	43.7	29.7	
%	Sol.		(93.2) _d (68.5) ^d		(96.2) (95.2)		(95.9)	(95.3)		(82.6)	(31.4)		(96.8)	(/6.5)		(95.7)	(96.4)		(66.4)	(98.4)		
	Sus.		93.8	l	95.0 95.2	1	96.1	96.7	ł	82.2	: 1		97.1	75.4		96.0	97.1	1	98.3	97.6		
	Sol. Sol., mg/l				977	789	685	603	7 60				989	927	909					659	647	
	Color Units		244	967	178 122	296	133	145	697	126	218		112	52	2	52	41	165	38	25	150	
	soluble		167	717	170 190	233	155	156	272	167	220		128	144	991	120	100	122	82	126	140	
Effluent (D)	BODS, mg/l Total Solu		188	767	168 197	304	172	180	977	208	248		144	188	141	131	120	143	114	134	163	
Effl	Sus. Sol., mg/l		(24) (133) ^d	(46)	(18) (19)	(171)	(17)	(16)	(35)	(15)	31)		(14) _d	(66)	(35)	(16)	6	(29)	(2)	9	(24)	
	Sus.		28	10	32 14	213	22	2 5	4 V	107	21		18	140	70	20	10	45	80	12	36	
	Hd		5.08	77.	3.92 4.89	7.41	5.21	4.69	7.40	5.18	7.90		4.91	5.23	74./	5.04	5.05	7.68	5.32	5.12	7.50	
	Sol. Sol., mg/1				705		669						632							703		
	Color Units		252	8	244 288	266	155	152	2	178	180	1	126	118	771	111	126	118	89	118	133	
()	Soluble		195 246	3	224 230	227	214	195	* 	242	747 747	1	186	186	8	168	148	158	107	194	25	
Influent (C)	BOD5, Total		332	arc	325 306	316	342	311	275	320	308	1	272	238		239	508	224	226	238	232	
Inf	Sus. Sol., mg/l		(337) (422)		(472) (210)		(415)	(428)		(408)	(300)		(437)	(422)		(374)	(248)		(309)	(371)		
	Sus. So mg/l		455 562		646 291	1	570	909	ţ	909	100		621	570	;	502	346	ļ	997	767	1	
	抵		7.35	1	7.48	1	8.42	7.79	1	7.75	6.		7.91	8.06	1	7.79	7.99	1	8,20	7.69	†	
	Composite		- 62	E;	1 7	×		7 3	Σ	(v Z			~ ;	E	1	2	Σ	e	2	Σ	
	Day		22		23		53			30			31			32			33			a,

Aglues in parentheses are volatile suspended solids.

Underscored values are based on average of IPC influent samples.

Composites

1 - 7.30 a.m. to 7:30 p.m. samples.

2 - 7:30 p.m. to 7:30 a.m. samples.

M - Mill composite analy2.4 in IPC lab.

Judge carry-over due to failure of sludge pump.

TABLE C-3. CLARIFIER STUDY AT COMBINED LOCKS - FERRIC CHLORIDE (75 mg/l Fe³⁺) PLUS NALCOLYTE 73C32 (0.75 mg/l)^{8,6}

	Color	50.0	i	-40.8 43.2	41.9	34.4
lon	D5 Soluble	20.6	-3.b	17.9 18.9 -11.0	20.3 25.5 5.8	17.3 21.4 7.0
% Reduction	BOD5 Total Sc	51.8	40.3	34.0 37.7 26.5	29.9 35.2 28.8	16.2 26.8 20.6
	Sus. Sol.	(99.0) (90.7)		(90.2) (86.7)	(88.1)	
		99.0	!	92.3	86.3	86.6
	Sol. Sol., mg/l	630	633	619		874
	Color Units	30	105	138 108 195	93 85 233	105 83 262
(a)	mg/l Soluble	100	144	87 193 191	212 213 260	196 220 240
Effluent (D)	BOD5, mg/l Total Solul	106	14.2	124 198 186	249 265 272	263 246 258
田	Sus. Sol. mg/l	33	(36)	(25) (48) (26)	(48) (64) (30)	
	Sus.	7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7	4	30 77 44	81 106 53	81 130 45
	Hd	5.54	8/.	6.30 6.57 7.89	6.70 6.80 7.85	6.75 5.15 7.48
	Sol. Sol., mg/l	591		593		819
	Color Units	09	1	98 190	160 163 	160
(၁)	mg/l Soluble	126	139	106 238 172	266 286 276	237 280 258
Influent (C)	BOD5, Total	220	877	188 318 253	355 409 382	314 336 325
In	Sol.,	(292)	Pump	(254) (361)	(403) (456)	
	Sus. Sol., mg/l	424 258	Failure of Sludge Pump	389	590 702	606 431
	Hd	8.10	re of	7.99 8.31	8.18	7.65
	Composite		n 1) 2) Failu M)		2 dd &	1 ਰ ਕ 2 ਨ
	Day	36	37	38	39	40

Avalues in parentheses are volatile suspended solids.
Underscored values are based on average of IPC influent samples.
Composites
1 7:30 a.m. to 7:30 p.m. samples.
2 7:30 a.m. to 7:30 a.m. samples.
M — Mill composite analyzed in IPC lab.
Heavy floc carry-over due to slade pump troubles.

Table c-4. Clarifier study at combined locks -- ferric sulfate (100 mg/l Fe $^{3+}$) 6

AUnderscored values are based on average of IPC influent samples. Composites
1 - 7:30 a.m. to 7:30 p.m. samples.
2 - 7:30 p.m. to 7:30 a.m. samples.
M - Mill composite analyzed by IPC lab.

TABLE C-5. CLARIFIER STUDY AT COMBINED LOCKS - ALUM (300 mg/l)^a

mg/1 Color Sus. Sol., mg/1 BODs, mg/1 Color Color 226 168 5.25 29 178 180 113 93 226 168 5.25 29 178 180 113 93 226 1520 103 202 190 117 97 234 113 4.98 249c 216 202 128 54 270 208 5.63 26c 244 226 228 94 243 113 4.98 249c 248 226 295 94 270 208 5.63 26c 244 226 236 94 243 1140 180 174 90 66 94 242 200 197 177 91 140 145 91 250 255 5.20 35 242 246 145 91 260 168				ū	Influent (()			Effl	Effluent (D)				% Rem	oval	
7.57 436 310 226 168 5.25 29 178 180 113 N.C. Sample Pump Failed 5.20 103 202 190 117 7.42 542 332 226 155 5.23 16c 198 195 117 7.12 544 332 226 155 5.23 16c 198 195 132 7.12 544 332 234 113 4.98 249c 244 226 295 7.53 488 336 270 208 5.63 266 244 226 296 7.59 437 317 242 246 246 230 348 7.54 419 336 213 215 10.18 140 180 177 7.62 296 256 24 246 246 346 7.86 283 222 166 253 10.00	O	omposite	ЬH	Sus. Sol., mg/l	BOD5, Total	-	Color Units	ЬH	Sus. Sol., mg/1	BODs, Total		Color	Sus. Sol.	BODs Total Solu	Ds. Solubles	Color
N.G. Sample Pump Failed 126 168 5.25 29 178 180 113 N.G. Sample Pump Failed 5.20 163 202 190 117 7.42 544 332 226 155 5.23 16c 198 195 117 7.12 544 332 226 155 5.23 16c 246 202 190 117 7.13 488 368 270 208 5.63 26c 244 226 153 7.53 488 368 270 208 5.63 26c 244 226 295 7.59 437 317 242 20 5.35 37 200 197 177 7.62 289 367 296 255 5.20 35 242 246 36 7.62 289 366 255 5.20 35 242 246 36 7.79<	1															
N.G. Sample Pump Failed 5.20 103 202 190 117 7.42 542 332 226 155 5.23 16 198 195 132 7.12 544 332 226 155 5.23 16 198 195 132 7.53 488 368 270 208 5.63 26 244 226 235 226 226 248 226 226 226 226 226 226 226 226 226 226 226 226 226 226 248 230 348 7.59 437 317 242 200 5.35 37 200 197 177 7.62 289 367 296 255 5.20 35 242 246 153 7.62 289 367 296 255 5.20 35 284 246 346 7.76 289 <			7.57	436			168	5.25	53	178	180	113	93.3	42.6	20.4	32.7
7,42 542 542 332 226 155 5.23 166 198 195 132 132 7,12 544 332 226 113 4.96 2496 216 202 118 7,53 488 368 270 208 5.63 266 244 226 295 7,59 437 317 242 200 5.35 37 200 197 177 7,75 419 336 213 215 10.18 140 180 174 90 7,62 289 367 296 255 5.20 35 242 246 153 7,62 289 367 296 255 5.20 35 242 246 153 7,62 289 268 202 168 4.61 246 246 146 36 7,86 283 268 202 168 4.61 246		ο;	X.G.	Sample		ed		5.20	103	202	961	117		1 1	! !), }
7.42 542 332 226 155 5.23 16c 198 195 132 132 7.12 544 332 234 113 4.98 249c 216 202 128 7.53 488 368 270 208 5.63 26c 244 226 295 7.59 437 317 242 200 5.35 37 200 197 177 7.62 289 367 296 255 5.20 35 242 246 153 7.62 289 367 296 255 5.20 35 242 246 153 7.62 289 367 296 255 5.20 35 242 246 153 7.62 289 268 202 168 4.61 24 140 38 7.79 293 268 202 168 4.61 24 164 145		E	!					67.7		287	27	577				ì
7.12 544 332 234 113 4.98 249 216 202 128 7.53 488 368 270 208 5.63 26 244 226 295 7.59 437 317 242 200 5.35 37 200 197 177 7.75 419 336 213 215 10.18 140 180 174 90 7.62 289 367 296 255 5.20 35 242 246 153 7.62 289 367 296 255 5.20 35 242 246 153 7.62 289 367 296 255 5.20 35 242 246 153 7.86 283 222 166 223 10.00 116 146 145 7.75 293 245 184 7.72 58 258 218 217		-	7.42	542	332	226	155	5,23	16	198	195	132	97.0	40.4	13.7	14.8
7.53 488 368 270 208 5.63 26 244 226 295 344 243 7.15 49 248 230 348 7.59 437 317 242 200 5.35 37 200 197 177 7.62 289 367 296 255 5.20 35 242 246 153 7.62 289 367 296 255 5.20 35 242 246 153 7.86 283 222 166 223 10.00 116 154 140 38 7.79 293 268 202 168 4.61 24 164 145 7.75 293 245 184 7.72 58 258 218 217 7.65 406 240 165 113 5.51 236 228 228		1 1	7.12	244	332	234	113	4.98	249	216	202	128	54.2ª	39.2	13.7	-13.3
7.59 437 317 242 260 5.35 37 200 197 177 7.59 437 317 242 200 5.35 37 200 197 177 7.75 419 336 213 215 10.18 140 180 174 90 7.62 289 367 296 255 5.20 35 242 246 153 7.86 283 222 166 223 10.00 116 154 140 38 7.79 293 268 202 168 4.61 24 165 145 7.65 406 240 165 17.72 58 2.58 218 217 7.80 181 281 250 168 6.20 26 26 228 105 7.80 181 281 250 168 6.20 26 258 228 228 228		2	7.53	488	368	270	208	5,63	2 6~	244	226	295	9.46	38.8	16.3	-41.8
7.59 437 317 242 200 5.35 37 200 197 177 7.62 289 367 296 255 5.20 35 242 246 153 7.62 289 367 296 255 5.20 35 242 246 153 7.86 283 222 166 223 10.00 116 154 140 38 7.79 293 268 202 168 4.61 24 163 164 145 7.75 293 245 184 7.72 58 258 218 217 7.65 406 240 165 113 5.51 22 204 200 52 7.80 181 281 250 168 10.15 147 258 228 105 7.80 181 284 217 7.51 47 222 216		×	ŀ	ì	344	243	1	7.15	49	248	230	348	;	27.9	5.3	1
7.75 419 336 213 215 10.18 140 180 174 90 7.62 289 367 296 255 5.20 35 242 246 153 7.86 283 222 166 223 10.00 116 154 140 38 7.79 293 268 202 168 4.61 24 163 164 145 7.65 406 240 165 113 5.51 25 258 218 217 7.80 181 281 250 168 10.15 147 258 105 7.80 181 281 250 168 10.15 147 258 228 105 7.80 181 281 250 168 10.15 147 258 228 105 7.80 181 284 217 7.51 47 222 216 73 <td></td> <td>41</td> <td>7.59</td> <td>437</td> <td>317</td> <td>242</td> <td>200</td> <td>5,35</td> <td>37</td> <td>200</td> <td>197</td> <td>177</td> <td>91.5</td> <td>36.9</td> <td>18.6</td> <td>11.5</td>		41	7.59	437	317	242	200	5,35	37	200	197	177	91.5	36.9	18.6	11.5
7.62 289 367 296 255 5.20 35 242 246 153 340 250 7.38 75 284 246 153 7.86 283 222 166 223 10.00 116 154 140 38 7.79 293 268 202 168 4.61 24 163 164 145 245 184 7.72 58 258 218 217 7.65 406 240 165 113 5.51 22 204 200 52 7.58 373 272 236 168 6.20 26 226 228 105 7.80 181 281 250 168 10.15 147 258 224 62 264 217 7.51 47 222 216 </td <td></td> <td>18</td> <td>7.75</td> <td>419</td> <td>336</td> <td>213</td> <td>215</td> <td>10,18</td> <td>140</td> <td>180</td> <td>174</td> <td>90</td> <td>9.99</td> <td>46.4</td> <td>18,3</td> <td>58.1</td>		18	7.75	419	336	213	215	10,18	140	180	174	90	9.99	46.4	18,3	58.1
7.86 283 222 166 223 10.00 116 154 140 38 7.79 293 222 166 223 10.00 116 154 140 38 7.79 293 268 202 168 4.61 24 163 164 145 7.65 406 240 165 113 5.51 22 204 200 52 7.58 373 272 236 168 6.20 26 236 228 105 7.80 181 281 250 168 10.15 147 258 224 62 - - 264 217 - 7.51 47 222 216 73		2	7.62	289	367	296	255	5.20	35	242	246	153	87.9	34.0	16.9	40.0
7.86 283 222 166 223 10.00 116 154 140 38 7.79 293 268 202 168 4.61 24 163 164 145 7.65 406 240 165 113 5.51 22 204 200 52 7.80 181 281 250 168 10.15 147 258 228 105 7.80 181 281 250 168 10.15 147 258 224 62 7.80 181 281 20 168 10.15 147 258 224 62 7.80 181 264 217 7.51 47 222 216 73		æ	1	ł	340	52	ł	7.38	75	284	246	342	Į	15.0	2.0	1
7.79 293 268 202 168 4.61 24 163 164 145 245 184 7.72 58 258 218 217 7.65 406 240 165 113 5.51 22 204 200 52 7.58 373 272 236 168 6.20 26 236 228 105 7.80 181 281 250 168 10.15 147 258 224 62 264 217 7.51 47 222 216 73		-	7.86	283	222	166	223	10,00	116	154	140	38	59.0	30.6	15.7	83.0
7,65 406 240 165 113 5,51 22 204 200 52 7,89 373 272 236 168 6.20 26 226 228 105 7,80 181 281 250 168 10.15 147 258 224 62 264 217 7.51 47 222 216 73		2	7.79	293	268	202	168	4.61	77	163	164	145	91.8	39.2	18.8	13.7
7,65 406 240 165 113 5.51 22 204 200 52 7,58 373 272 236 168 6.20 26 236 228 105 7,80 181 281 250 168 10.15 147 258 224 62 264 217 7.51 47 222 216 73		×	ł	1	245	2	ł	7.72	28	258	218	217	!	-5.0	-18.5	1
7.58 373 272 236 168 6.20 26 236 228 105 7.80 181 281 250 168 10.15 147 258 224 62 <u>264</u> <u>217</u> 7.51 47 222 216 73		1	7.65	406	240	165	113	5.51	22	707	200	52	94.6	15.0	-21.2	54.0
7.80 181 281 250 168 10.15 147 258 224 62 <u>264 217</u> 7.51 47 222 216 73		7	7.58	373	272	236	168	6.20	56	236	228	105	93.0	13.2	3.4	37.5
<u>264</u> <u>217</u> 7.51 47 222 216 73		7Y	7.80	181	281	250	168	10.15	147	258	224	62	18.8	8.2	10.4	63.1
		×	1	}	564	217	ŀ	7.51	47	222	216	73	1	15.9	0.5	1

**Underscored values are based on average of IPC influent samples.

Composites

1 - 7:30 a.m. to 7:30 p.m. samples

2 - 7:30 p.m. to 7:30 a.m. samples

M - mill composite analyzed by IPC lab.

Sheavy floc carry-over due to sludge pump troubles.

TABLE C-6. CLARIFIER STUDY AT COMBINED LOCKS WITH AND WITHOUT ALUM (300 mg/l) PLUS HERCOFLOC 812.3 (0.75 mg/l)

			Inf	Influent (C	~			Rff		7			% Reduction	tion	
Day	Composite	版	Sus. Sol., mg/l	BOD ₅ , mg/ Total So	mg/l Soluble	Color Units	рН	Sus. Sol., mg/l	BOD5, Total	mg/l Soluble	Color Units	Sus. Sol,	BO Total	BOD ₅ al Soluble	Color
57	100	7.76	997	342	256	136	6.11	86 86	252	234	120 156	91.8	26.3	8.6	11.8
	ıΣ	1	: 1	369	275	1	7.45	84	307	276	238	1	16.8	0	ł
28	1 1A	7.49	670 495	308 386	188 234	184 172	6.01 6.12	54 118	185 239	180 216	164 200	91.9	39.9 38.1	4.2	10.9
	7 ×	7.52	561	320	210	207	7.29	63 56	242 236	218 218	246 254	88.8	24.4 30.2	-3.8	-18.8
	Alum pump failed	j failed													
11	3 0 ⊢	8.31	1164 431	553 377 46 <u>7</u>	362 276 319	239 262 	7.05 5.21 7.81	187 130 56	383 236 320	310 228 279	101 74 254	83.9 69.8	30.7 37.4 31.5	14.4 17.4 12.5	57.7
72	Z 2 1	8.60	2340 471 	527 372 450	347 200 274	234 140 	6.50 7.05 7.80	70 91 76	260 230 304	239 160 260	106 74 212	97.0 80.7 	50.7 38.2 32.4	31.1 20.0 5.1	54.7 47.1
73	1 2 W	8.20	527 591 	272 316 294	156 115 136	117	7.08 7.20 7.58	58 90 44	122 170 145	114 141 131	82 98 188	84.8	55.1 46.2 50.7	26.9 -22.6 3.7	29.9

**Bunderscored values are based on average of IPC influent samples. Composites

1 - 7:30 a.m. to 7:30 p.m. samples.

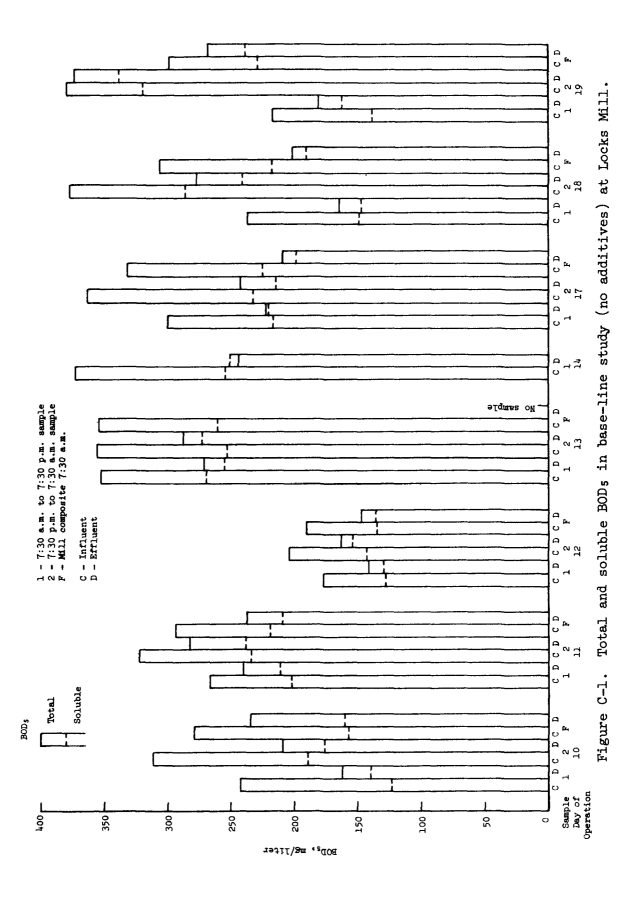
2 - 7:30 p.m. to 7:30 a.m. samples.

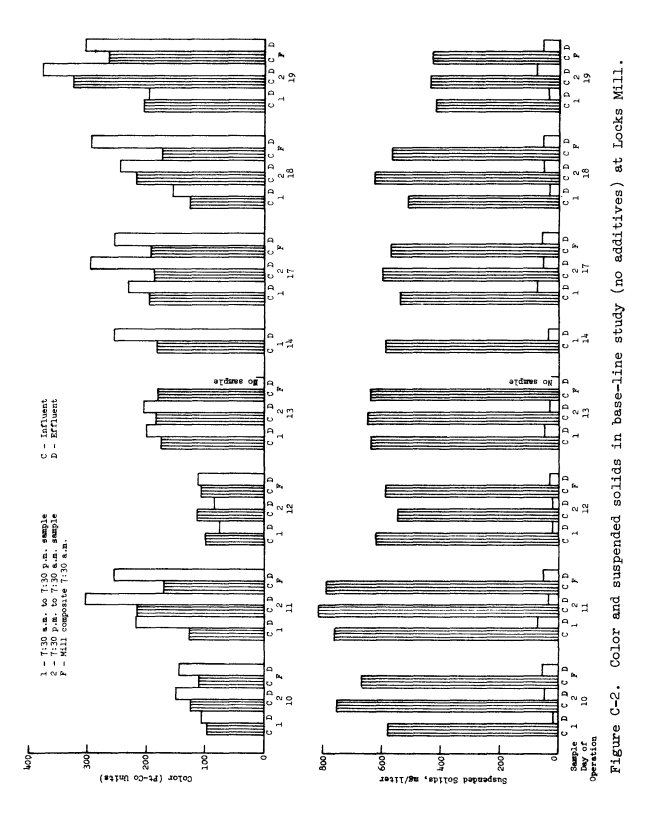
M - Mill composite analyzed by IPC lab.

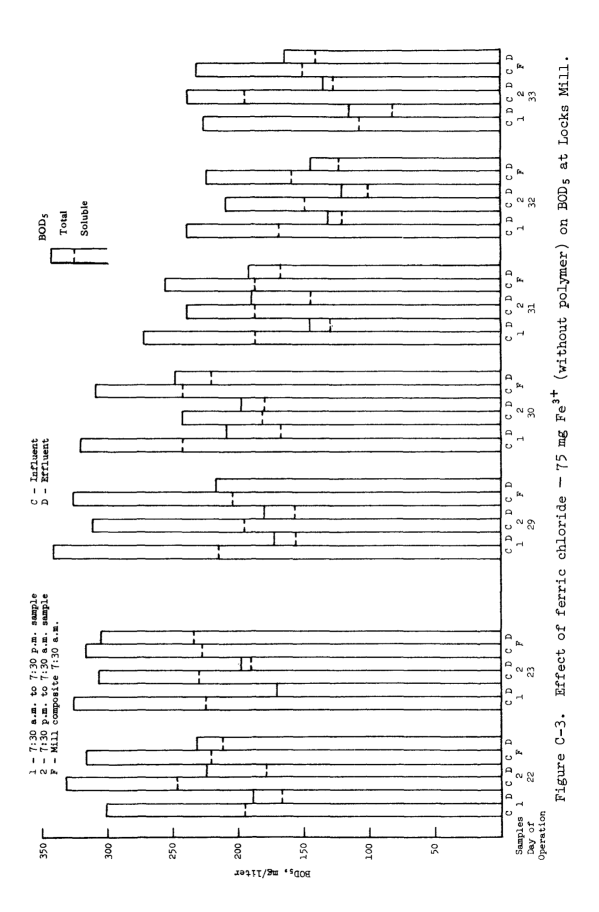
TABLE C-7. CLARIFIER STUDY AT COMBINED LOCKS - SECOND BASE LINE (GRAVITY SEDIMENTATION AFTER 0.010-INCH SIEVE) WITHOUT ADDITIVES

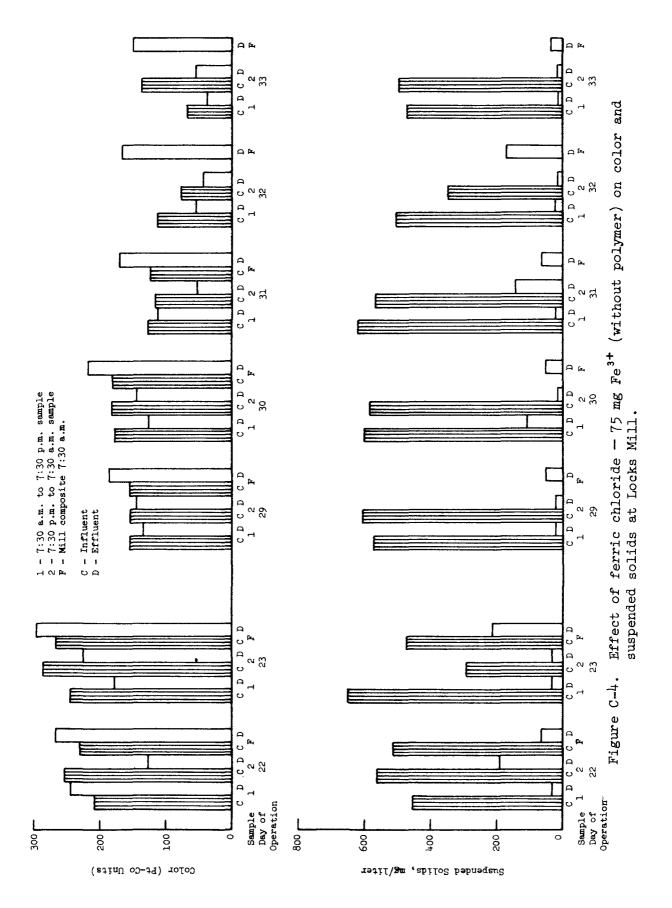
Soluble Units PH mg/1 Total Soluble Units Sus. Sol. Total		ı		Inf Bus. Sol	Influent (C)		Color		Ef Sus. Sol.)) mg/1	Color		% Reduction BODs	Juction BODs	
157 188 7.68 32 154 156 242 93.0 37.4 0.6 406 500 7.78 52 514 450 656 92.7 9.8 -10.8 282 - 7.45 50 312 292 326 92.7 9.8 -10.8 228 - 7.45 48 234 191 340 91.3 15.8 12.8 224 - 7.63 48 262 200 208 12.1 10.7 224 - 7.63 48 262 200 208 86.7 31.1 10.7 224 - 8.35 69 206 200 208 86.7 33.1 -4.2 228 - 8.02 20 20 20 86.7 33.1 -4.2 204 160 9.12 47 222 200 192 34.8	Composite pH mg/l		mg/1		Total	Soluble	Units	Hd	mg/1	Total	Soluble	Units	Sus. Sol.	Total	Soluble	Color
288	1 7.52 461		461		246	157	188	7.68	32	154	156	242	93.0	37.4	9.0	-28.7
228 238 7.85 57 243 220 328 90.0 23.3 3.5 219 212 7.87 48 234 191 340 91.3 15.8 12.8 224 7.63 48 262 200 288 12.1 10.7 192 152 8.35 69 206 200 200 86.7 33.1 -4.2 264 204 8.42 60 302 275 278 88.5 26.3 -4.2 208 206 207 202 202 202 203 34.8 -1.8 206 8.73 47 208 202 200 90.7 36.2 1.0 206 8.73 44 276 246 230 90.7 36.2 1.0 206 8.88 44 276 246 230 90.8 2.0 2.0			3		804	40 6	<u> </u>	7.45	25 20	312	450 292	392	7.76	23.5	-10.8	-31.2
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Aunderscored values are based on average of IPC influent samples. Composites
1 - 7:30 a.m. to 7:30 p.m. samples
2 - 7:30 p.m. to 7:30 a.m. samples
M - Mill composite analyzed by IPC lab.









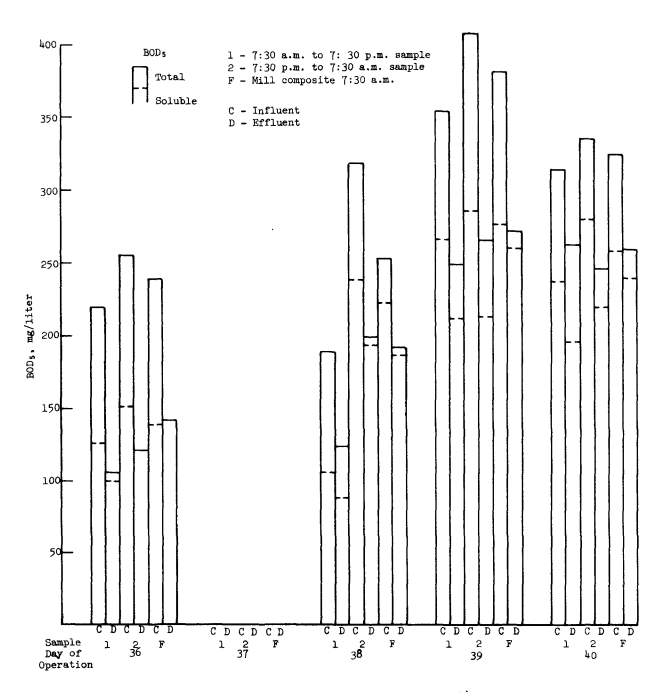
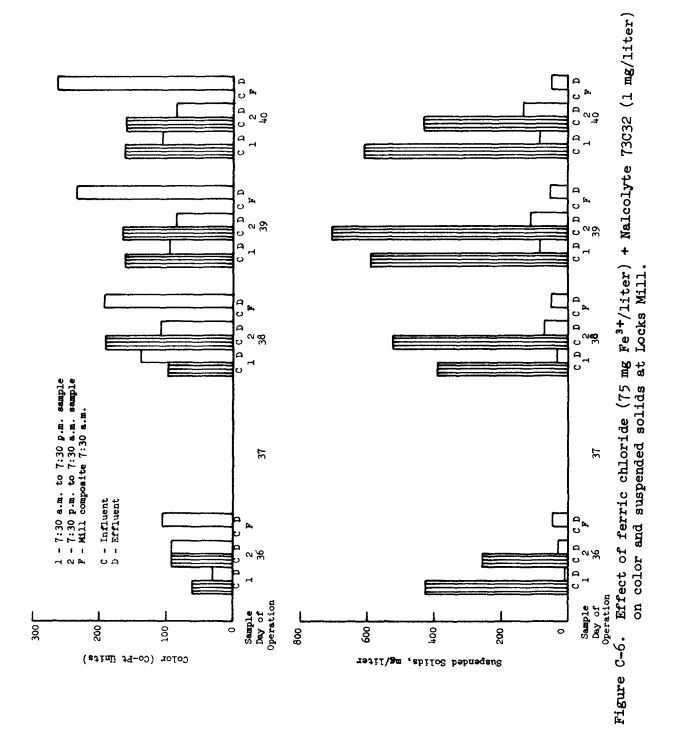


Figure C-5. Effect of ferric chloride (75 mg $Fe^{3+}/liter$) + Nalcolyte 73C32 (1 mg/liter) on BOD₅ at Locks Mill.



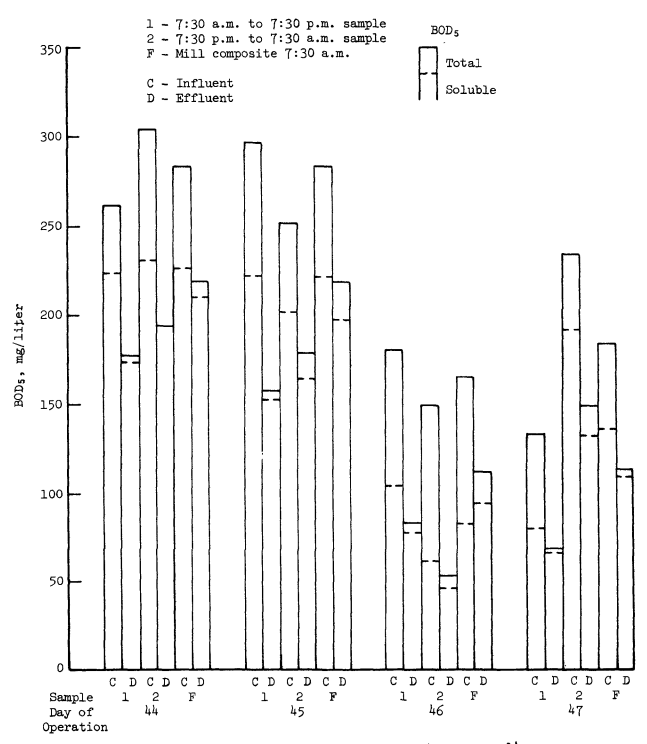


Figure C-7. Effect of ferric sulfate (100 mg ${\rm Fe}^{3+}/{\rm liter}$) on ${\rm BOD}_5$ at Locks Mill

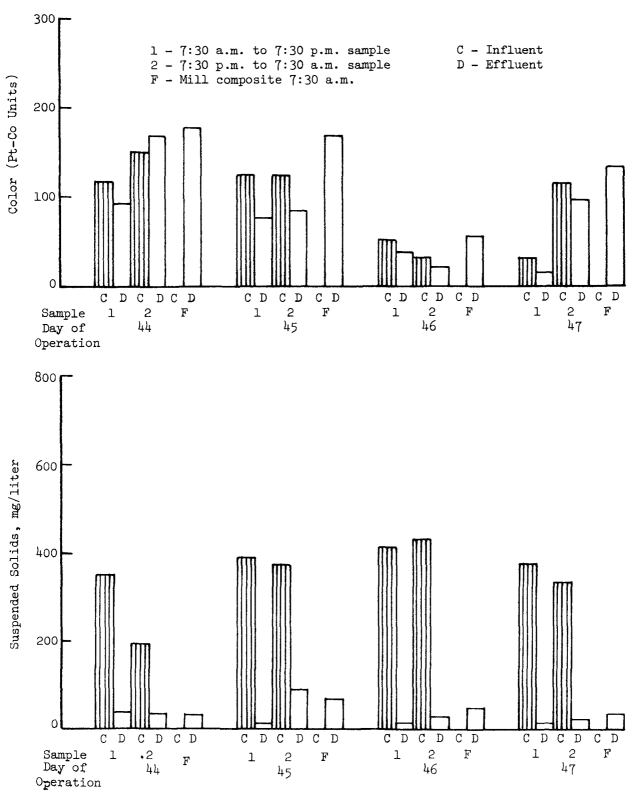
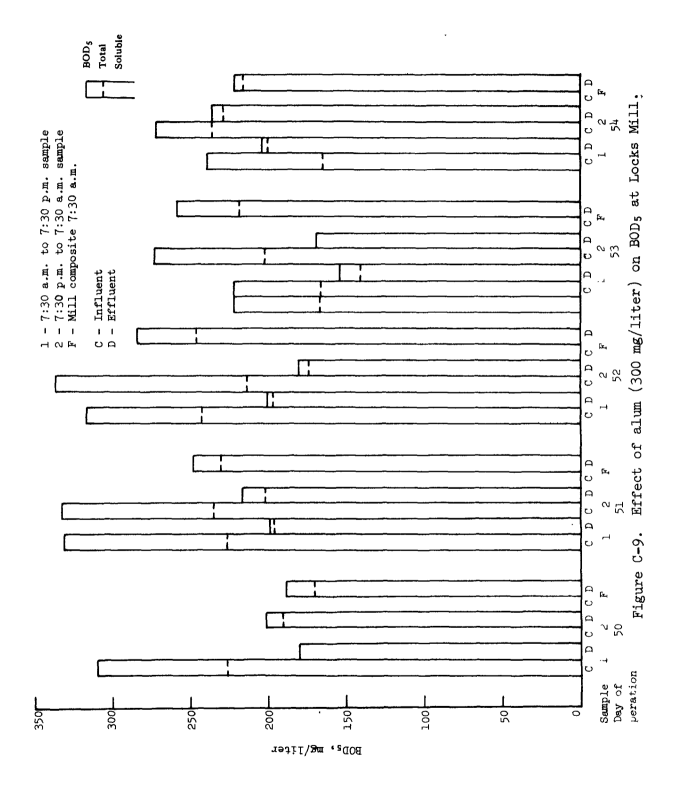


Figure C-8. Effect of ferric sulfate (100 mg Fe³⁺/liter) on color and suspended solids at Locks Mill.



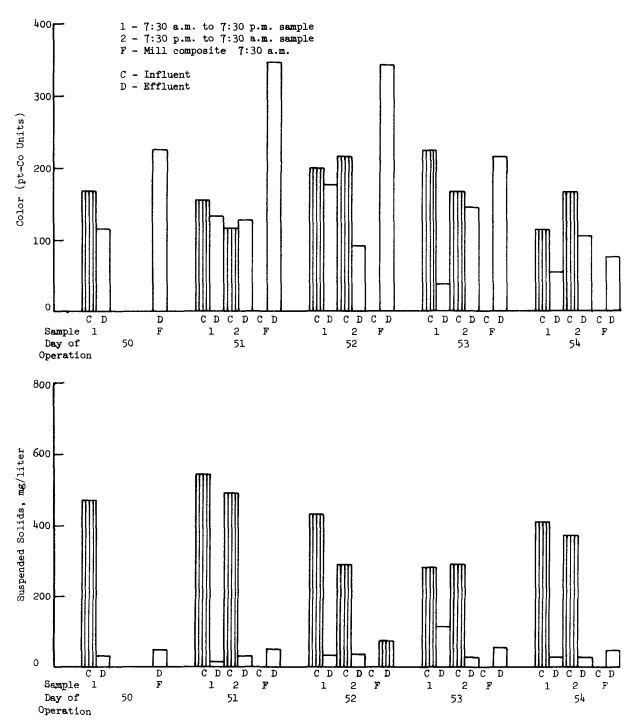


Figure C-10. Effect of alum (300 mg/liter) on color and suspended solids at Locks Mill.

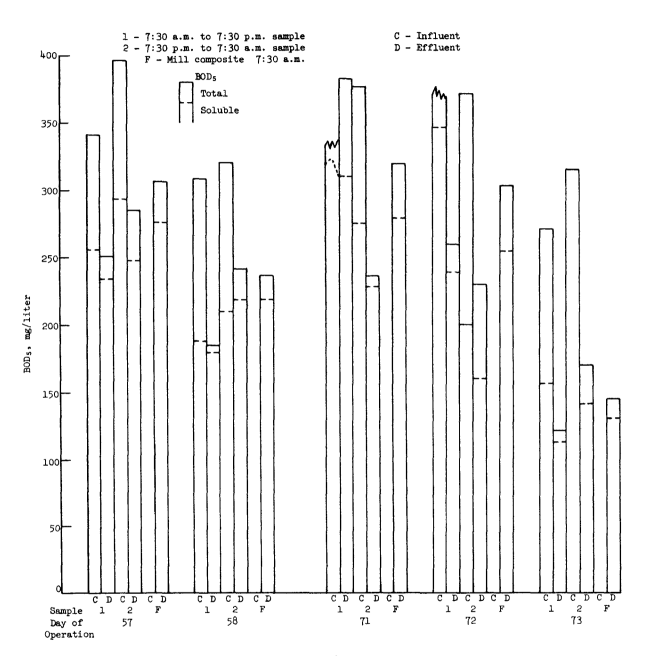
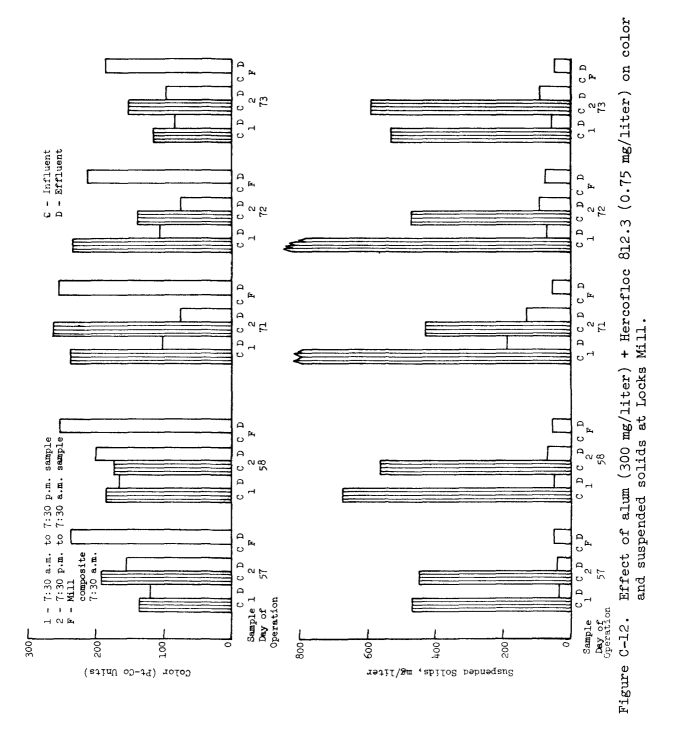
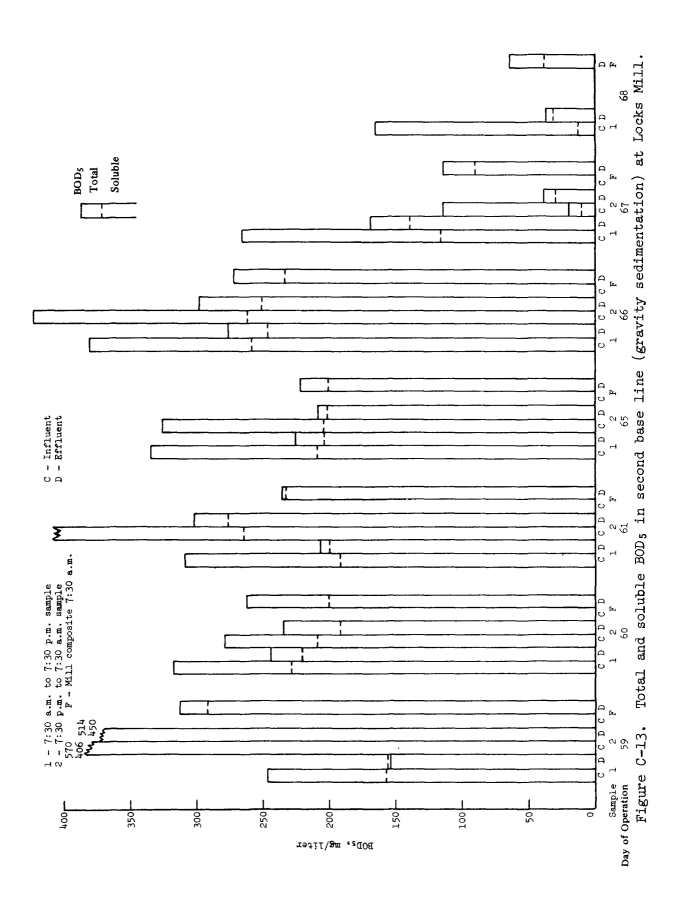
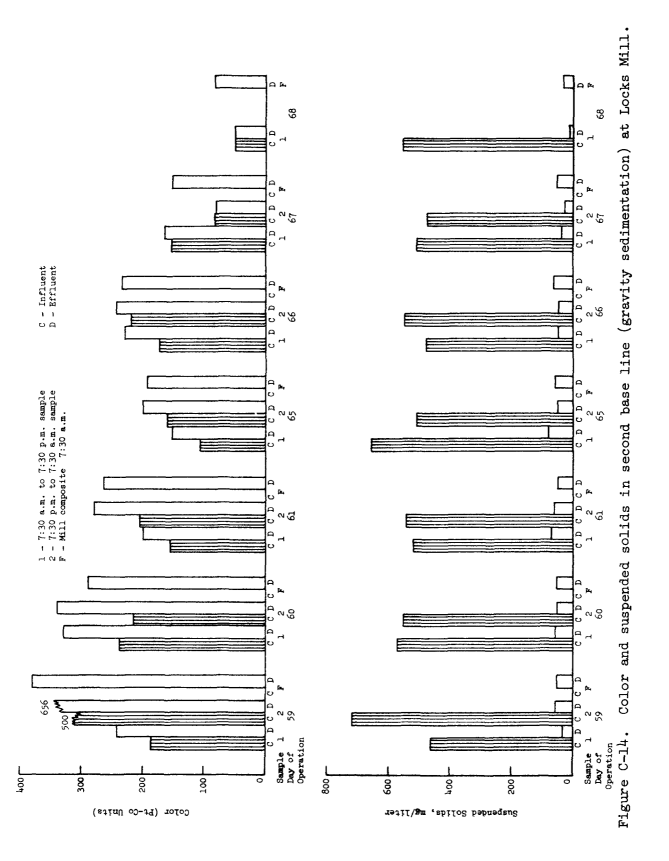


Figure C-ll. Effect of alum (300 mg/liter) + Hercofloc 812.3 (0.75 mg/liter) on BOD₅ at Locks Mill.







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15. SUPPLEMENTARY NOTES

16. ABSTRACT

This project was directed to evaluating means for increasing BOD removal from primary treatment systems treating pulp and paper wastes. An improved understanding of the optimal conditions for soluble and colloidal BOD removal should permit increasing efficiency in total organic removal.

The initial phase surveyed 12 mills to obtain data on total and soluble BOD5, COD, suspended solids and color removal from sedimentation systems. This data was used to select mill effluents for additional study. These laboratory studies showed that, with proper flocculating agents, soluble BOD removal could be markedly increased. Soluble BOD was defined as that organic load passing through a 0.45 µm filter.

Measurement of sludge volumes and sludge dewatering characteristics produced from use of iron salts, alum and polymers was outside the objective of the study.

Gel chromatography studies showed that low molecular weight biodegradable residues and colloidal materials were flocculated and removed. Studies with model compounds indicated that increased removal is apparently related to pH of the solution and to functional groups, chain length, branching and solubility of the compound.

Chemical costs may range from 3ϕ to 10ϕ per 1000 gallons. Cost reduction and improved clarifier performance can be achieved by elimination of overflows and spills within the mill that are antagonistic to efficient sedimentation due to dispersant action.

17.	KEY WORDS AND DOCUMENT ANALYSIS
a. DESCRIPTORS	b.IDENTIFIERS/OPEN ENDED TERMS C. COSATI Field/Group
Precipitation (chemistry) Coagulation Flocculating Oxygen demand	Primary treatment Water pollution control Pulp waste treatment 13B Paper waste treatment Chemical degradation
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