

Institute of Paper Science and Technology Atlanta, Georgia

IPST Technical Paper Series Number 638

Low Effluent Processing in the Pulp and Paper Industry: Electrodialysis for Continuous Selective Chloride Removal

P.H. Pfromm

February 1997

Submitted to Separation Science and Technology

INSTITUTE OF PAPER SCIENCE AND TECHNOLOGY PURPOSE AND MISSIONS

The Institute of Paper Science and Technology is a unique organization whose charitable, educational, and scientific purpose evolves from the singular relationship between the Institute and the pulp and paper industry which has existed since 1929. The purpose of the Institute is fulfilled through three missions, which are:

- to provide high quality students with a multidisciplinary graduate educational experience which is of the highest standard of excellence recognized by the national academic community and which enables them to perform to their maximum potential in a society with a technological base; and
- to sustain an international position of leadership in dynamic scientific research which is participated in by both students and faculty and which is focused on areas of significance to the pulp and paper industry; and
- to contribute to the economic and technical well-being of the nation through innovative educational, informational, and technical services.

ACCREDITATION

The Institute of Paper Science and Technology is accredited by the Commission on Colleges of the Southern Association of Colleges and Schools to award the Master of Science and Doctor of Philosophy degrees.

NOTICE AND DISCLAIMER --

The Institute of Paper Science and Technology (IPST) has provided a high standard of professional service and has put forth its best efforts within the time and funds available for this project. The information and conclusions are advisory and are intended only for internal use by any company who may receive this report. Each company must decide for itself the best approach to solving any problems it may have and how, or whether, this reported information should be considered in its approach.

IPST does not recommend particular products, procedures, materials, or service. These are included only in the interest of completeness within a laboratory context and budgetary constraint. Actual products, procedures, materials, and services used may differ and are peculiar to the operations of each company.

In no event shall IPST or its employees and agents have any obligation or liability for damages including, but not limited to, consequential damages arising out of or in connection with any company's use of or inability to use the reported information. IPST provides no warranty or guaranty of results.

The Institute of Paper Science and Technology assures equal opportunity to all qualified persons without regard to race, color, religion, sex, national origin, age, disability, marital status, or Vietnam era veterans status in the admission to, participation in, treatment of, or employment in the programs and activities which the Institute operates.

Low Effluent Processing in the Pulp and Paper Industry:

Electrodialysis for Continuous Selective Chloride Removal

Peter H. Pfromm
Institute of Paper Science and Technology
Atlanta, GA

To be submitted to Separation Science and Technology

Abstract

Pollution prevention is currently a major focus of the U.S. pulp and paper industry. Significant process changes are inevitable to implement low effluent processing. The kraft pulping process is prevalent for the production of wood pulp. About 50 million tons of wood pulp are produced annually in the United States alone using the kraft process. Water consumption is currently roughly between 30 and 200 m³ of water per ton of air dry bleached kraft pulp. In-process recycling of water is now being implemented by many mills to reduce the use of increasingly scarce water resources and to reduce the need for wastewater treatment.

Mass balance considerations and industrial experience show that non-process elements (NPE's), which are detrimental to the kraft process, such as chloride and potassium, will quickly build up once water use is significantly reduced. High concentrations of chloride and potassium can cause corrosion and lead to more frequent mill shutdowns due to fouling of heat exchanger surfaces in the kraft recovery furnace. Electrodialysis with monovalent selective anion and cation exchange membranes was explored here to selectively remove chlorine as sodium and potassium chloride from a feed stream with very high ionic strength. Experiments with model solutions and extended tests with the actual pulp mill materials were performed. Very good selectivities and current efficiencies were observed for chloride over sulfate. The outstanding performance of the process with actual mill materials containing organic and inorganic contamination shows great promise for rapid transfer to the pilot scale. This work is an example of the usefulness of membrane separations as a kidney in low effluent industrial processing.

Keywords: Electrodialysis, monovalent selective membranes, pulp and paper, kraft process, pollution prevention, capital and operating costs

Introduction and Background

The theory and practice of the kraft pulping process is described in detail elsewhere (Grace et al., 1991). Wood is exposed to an aqueous solution of sodium sulfide and sodium hydroxide at elevated temperature and pressure in a batch or continuous reactor. Thereby, the lignin is solubilized and the cellulose fibers are liberated. The solution of lignin and spent inorganic chemicals is then separated from the cellulose fibers. This solution is concentrated by evaporation and used as fuel in the kraft recovery furnace. In this furnace, the lignin is burned to produce steam for the mill, and the inorganic chemicals are recovered as a molten salt mixture of sodium carbonate and sodium sulfide. This salt mixture is then dissolved in water and reprocessed to recover and reuse the active pulping chemicals.

One of the challenges for the pulp and paper industry is low effluent processing to minimize environmental impact and water use. A serious impediment to low effluent processing is the enrichment of certain elements, termed non-process elements (NPE's) in the process fluids when recycling of water is implemented. The term NPE loosely describes materials, which are not needed in the process, but which are introduced through raw materials and replacement chemicals. The NPE's chlorine and potassium are attracting much attention due to their negative impact on the kraft pulping cycle. The good solubility of chloride and potassium ions in alkaline solutions is the cause of this accumulation.

Corrosion increases with the chloride concentration in kraft pulping fluids (Ahlers et al., 1976). Chloride and potassium also significantly increase shut downs of kraft recovery furnaces by fouling of heat exchanger surfaces with sticky deposits (Reeve et al., 1981; Tran, 1986). The kraft recovery furnace is crucial to pulp mill operation. It produces most of the steam and energy used in a pulp mill by burning the unwanted organic wood components. Shutdowns of the recovery furnace cause costly pulp mill outages.

Chloride contributes to the inorganic deadload in the kraft pulping process. This can reduce the actual firing capacity of a recovery furnace for organic material if the furnace permit is written for overall

solids fired. This can limit mill production. Even fractions of a percent of capacity decrease are economically very significant in a commodity industry such as pulp and paper.

Input of chloride into the pulping process can be minimized, but not eliminated, due to its presence in wood (Keitaanniemi and Virkola, 1982; NCASI, 1994). Chloride is highly soluble under the conditions encountered in kraft pulping and bleaching. If low effluent operation is pursued, tightening of the liquor cycles will lead to increasing concentrations of the soluble NPE's. A highly specific purge is therefore needed for chloride (Karjalainen et al., 1972; Moy et al., 1974).

The introduction of bleach plant effluents into the kraft recovery cycle is widely regarded as a way to low effluent bleached kraft pulp production. The basic idea is to burn unwanted organics in the recovery furnace, instead of discharging them with water. However, together with the organic material, inorganic matter is recycled. This further emphasizes the need for a specific purge of soluble inorganics (Rapson, 1967; Reeve, 1976). This is especially true if chlorine is present in some form in the bleaching chemicals (Holman et al., 1994).

A significant number of processes for removal of chloride from the kraft pulping process are described in the literature and in patents. The good solubility in kraft pulping liquors makes chloride difficult to purge without loss of large amounts of valuable chemicals that are also present. Purging of chloride as hydrochloric acid with the kraft recovery furnace flue gas has been considered, but does not appear to be a desirable long-term option because this would constitute only a shift of the emission problem. An economical chloride purge could be achieved by a highly selective removal process, or by discarding part or all of a chloride-rich process stream.

Capital investment, operating cost, maintenance, ease of operation, environmental impact, and the regulatory situation will all impact the choice for an optimum solution. The best choice will certainly vary from mill to mill.

An advantageous point for a selective chloride purge is the electrostatic precipitator that removes dust from the flue gas of the kraft recovery furnace. Several weight % of the total fired solids in the furnace are recovered as electrostatic precipitator dust (ESP dust) from the flue gas. This dust is then recycled to

the process. It has been found that chloride and potassium are enriched in the ESP dust, when compared with their relative concentrations in the main process stream (black liquor) (Tran, 1986). This makes ESP dust a good candidate for a chloride purge process.

Differential solubility in water is clearly the selective chloride removal mechanism that has attracted the most interest in the past (for example, Moy et al., 1974; Wright et al., 1956). Excellent selectivity is inherent in crystallization processes, but a subsequent liquid/solid separation is necessary. This will cause some carryover of chloride-containing liquid. The chloride purge stream, if dissolved ESP dust is treated, will contain organic materials. These organics stem from not completely combusted organics in the fuel (black liquor). The capital investment and maintenance for a crystallization/filtration process are not insignificant. Crystallizer performance, handling of two-phase systems, and startup/shutdown procedures may be other concerns.

The selective electrodialytical removal of chloride from dissolved ESP dust with chloride selective membranes is reported here. A similar technology involving a leaching step and subsequent electrodialysis has been reported (Maruko et al., 1980). The work reported here differs in that no leaching or pretreatment of ESP dust was needed for successful long-term chloride separation experiments with actual mill materials. A schematic diagram of the process is shown in Figure 1.

It is important to note that the goal is not to reduce the chloride concentration in the ESP dust to near zero, but to continuously remove the chloride input to the system to maintain steady state. This is a very advantageous situation where a membrane process serves as a kidney, rather than an ultimate polishing process. The result of the work presented here is a simple process for retrofitting of existing mills. Estimates of operating costs and capital investment costs are given below.

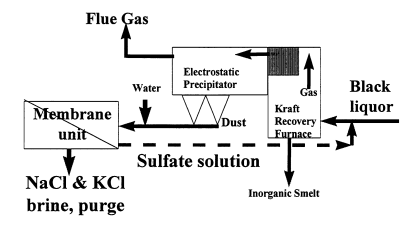


Figure 1: Schematic diagram for selective chloride purge from dissolved ESP dust in a kraft pulp mill.

Selective electrodialysis of chloride/sulfate solutions has been investigated elsewhere at concentrations far lower than reported here (Saracco et al., 1993). In electrodialysis, the selectivity at high ionic strengths can usually be expected to decline. The results shown here give an example of good performance of commercial monovalent selective electrodialysis membranes, even at high ionic strengths.

Experimental

Materials

Mill materials

In the experiments reported here, ESP dust from a kraft mill in the Southeastern United States was used. The dust is a white powder (micron-size particles, free flowing). It dissolves easily in water, forming a brownish solution, where the colored organics settle over night, leaving a clear supernatant. The ability to settle under gravity shows the rather large particle size of the organic material. No settling was allowed for any of the electrodialysis experiments. The composition of the dust is reported in Table 1. The organic carbon is either a product of incomplete combustion or direct carryover of lignin from the black liquor fuel fed to the kraft recovery furnace.

Table 1: Main components of dissolved electrostatic precipitator dust from a southeastern kraft pulp mill.

Cl ⁻	CO ₃ ***	SO ₄ **	Na^+	K^{+}	Org. Carbon		
weight%							
3.46	2.35	57.15	25.09	10.74	0.10		

Mismatch of electrochemical equivalents

anions/cations:

0.03%

Mass balance, total wt% accounted for:

98.89%

For the experiments with mill materials, a solution with 143 grams of ESP dust per liter of mill water was made and used as the diluate.

Components such as calcium, metals, and transition metals were not analyzed here. Both the organic and inorganic composition of ESP dust will vary from mill to mill, and with time. Inorganic mill inputs vary due to geography (water supply, wood supply), seasonal changes of the wood, and impurities from replacement chemicals. Organic components in the ESP dust may vary with the firing strategy of the recovery furnace. Mill process water was used to dissolve the ESP dust for the chloride removal experiments. Main anions and cations in the mill process water were calcium (31 ppm), carbonate (93 ppm), and sulfate (28 ppm).

Model solutions

A representative composition for the model diluate (simulated ESP dust) was chosen based on the literature and on samples from several mills. The solution concentration was chosen to not exceed any solubility limits. Table 2 shows the initial diluate, concentrate, and electrode rinse compositions.

Table 2: Initial composition of model diluate (simulated ESP dust), concentrate, and electrode rinse solutions (all in deionized water).

	SO ₄ **	Cl ⁻	CO ₃ **	K^{+}	Na ⁺	
Diluate (Simulated ESP dust)	81.39	7.75	[g/l] 5.61	8.89	43.05	
Concentrate	4.02	-	-	-	1.93	
Electrode rinse	4.02	-	-	-	1.93	

The compositions for the electrode rinse solution and the initial concentrate solution were identical for all experiments, regardless of the diluate. The diluate composition was either as reported above or it was actual ESP dust from the mill dissolved in mill process water.

Membranes

The membranes were donated for this work by Tokuyama Soda Co., LTD. The membranes were NEOSEPTA® CMS (monovalent selective cation exchange membrane) and NEOSEPTA® AMS (monovalent selective anion exchange membrane). Two standard cation exchange membranes (NEOSEPTA® CM-1) were used for the ends of the stack. The bulk of AMS and CMS membranes consists of a strongly acidic (cation exchange) and strongly basic (anion exchange) resin, respectively. Both membranes contain a reinforcing fabric and are highly selective for monovalent ions, as measured in sea water and reported in the manufacturer's literature.

Inorganic salts

The inorganic salts used here were technical grade for sodium sulfate and analytical grade for sodium chloride, sodium carbonate, and potassium sulfate.

Water

Deionized water was used for all solutions. The exception is the diluate solution for experiments with actual mill ESP dust. The ESP dust was dissolved in mill process water (see Mill Materials above).

Electrodialysis System and Experimental Details

A laboratory electrodialysis system manufactured by Stantech, Inc. (Hamburg, Germany) was used. This system has an active cross-sectional membrane area of 100 cm² (10 cm by 10 cm). An

electrodialysis unit consists of a parallel arrangement of alternating diluate and concentrate cells, separated by anion and cation membranes. The setup is similar to a conventional filter press. A polymeric netting is placed inside of the frames forming the cells to minimize boundary layer effects.

Four diluate and five concentrate cells were used for the experiments. The effective cell thickness (distance from membrane to membrane) is 0.25 cm. The superficial average flow velocities were 4.4 cm/s in the diluate cells and 4.9 cm/s in the concentrate cell. The electrodes were rinsed in series from anode to cathode with a flow of 80 l/hr. The stack configuration and the goal of the separation process are shown in Figure 2. The outermost cation exchange membranes of the stack were standard membranes. All other anion and cation exchange membranes were monovalent ion selective.

In Figure 2, a repeat unit is indicated. Up to several hundred of these repeat units are used in commercial systems. The goal of the process investigated here is to selectively remove chloride as sodium and potassium chloride while retaining the sulfate, which is recycled to the kraft recovery furnace.

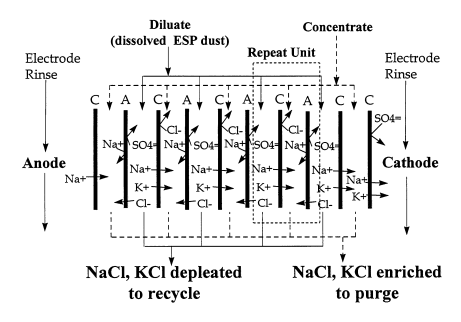


Figure 2: Laboratory stack setup for chloride removal from dissolved ESP dust. (A, C: monovalent selective anion and cation exchange membranes, respectively)

The experiments were run in batch mode with samples of 140 milliliters drawn at time intervals. A total electrical current of one ampere was passed through the stack, which corresponds to a current density of 10

mA/cm². Limiting current density experiments showed that this is far below the critical current density where boundary layer effects start to become important.

In batch experiments for electrodialysis, the concentrate circuit must be started with some electrical conductivity so that the electrical resistance of the stack is initially not too high. High electrical resistance would otherwise cause effects such as water splitting at the ion depleted membrane boundary layers. In our runs, sodium sulfate was used to supply this initial conductivity. In a full-scale unit, the concentrate conductivity would be supplied by a small recycle stream from the concentrate outlet (purge) to the concentrate inlet of the stack.

As indicated individually for each run reported here, 4 to 5.86 liters of diluate (actual ESP dust dissolved in mill process water, Table 1, or model ESP dust solution, Table 2), 4 to 5.86 liters of concentrate (see Table 2), and 4 liters of electrode solution (see Table 2) were used. The solution temperatures rose slightly during each experiment from room temperature (20°C) to about 27°C at maximum, mainly through heat transfer from the centrifugal pumps.

Analytical

The cations sodium and potassium were analyzed by atomic absorption spectroscopy. The anions carbonate, sulfate, and chloride where analyzed by ion chromatography.

Results and Discussion

Model solutions: chloride selectivity

Figure 3 shows the results from electrodialysis of a model solution to simulate ESP dust (initial composition, see Table 1). Figure 3 shows the electrochemical equivalents of chloride and sulfate transported from the diluate to the concentrate. Also shown is the theoretical maximum of transported ions, as calculated by integrating the total electrical current that passed through the unit multiplied by the number of diluate cells.

The sulfate and carbonate concentrations remain constant within the accuracy of the analysis. The chloride concentration in the concentrate increases rapidly, which shows the selective transport of chloride.

At the end point of this run, 65.9% of the chloride had been transferred from the diluate to the concentrate. The overall average current efficiency for chloride removal, when comparing the electrochemical equivalents of chloride transported with the total electrical current, was 73%.

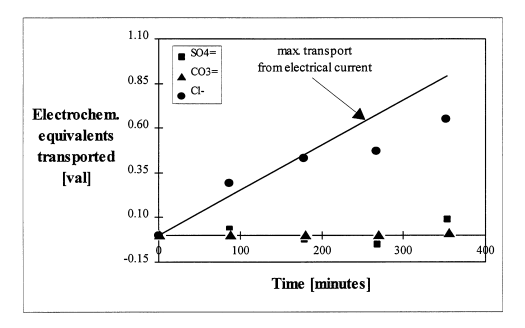


Figure 3: Ion transport by batch electrodialysis of model ESP dust solution (20.8-26.2 °C; 10 mA/cm²; diluate, concentrate: 4 liters each).

It is clear that even with the highly concentrated feed solution containing a very large excess of divalent sulfate ions over chloride ions the membranes are still very selective for monovalent anions. This encouraging result led us to initiate tests with the actual materials from a pulp mill.

Long-term tests with mill materials

Based on the encouraging selectivity for chloride found with the model solutions, long-term tests were initiated with mill materials. Figure 4 shows results from the first test with mill materials. Then, a number of standard separation runs with mill materials were performed with the same set of membranes (these runs are not shown). Also shown in Figure 4 are data for an experiment after a combined 86 hours of testing of this same set of membranes with the mill materials.

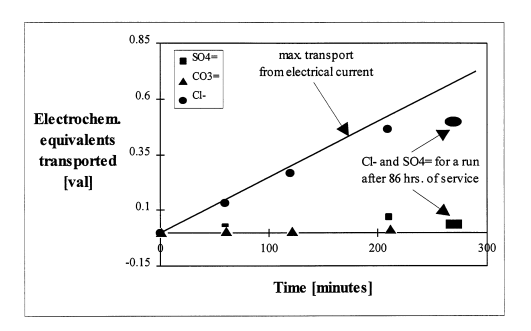


Figure 4: Ion transport by batch electrodialysis of actual ESP dust dissolved in mill water. Dechlorination performance of the same set of membranes before and after 86 hours of service with actual mill ESP dust solution (20.2-24.6 °C; 10 mA/cm²; 5.86 l diluate, 5.86 l concentrate first run; 4.86 l diluate, 4.86 l concentrate final run after 86 hours).

When comparing the initial run with the run after 86 hours of experiments with mill materials, no significant deterioration is detected. No special chemical or physical cleaning procedures were performed on the membranes during this extended series of tests. For storage between batch experiments, the stack was disassembled and the membranes sprayed with 0.5 molar sodium chloride solution from a laboratory wash bottle. The membranes were then stored in 0.5 molar sodium chloride solution. This was done four times during the time of performing the 86 hours of runs with actual materials. The loose brown deposits found on the diluate side of the membranes washed off quite easily, indicating that a simple backflush will likely suffice for in-place cleaning of a full-scale application.

Figure 5 shows the stack voltage as measured between the anode and cathode as a function of batch run time for the two runs shown in Figure 4. It is clear that the electrical resistance is not increased by extended runs with the actual mill materials. Therefore, fouling appears not to be a serious problem in this process. Dissolved anionic organic molecules such as sulfonated lignin could potentially foul the membranes. The high ionic strength of the feed solution may advantageously contribute to precipitation of any potentially soluble organics in the ESP dust.

The initial decrease of the stack voltage is due to improved conductivity of the concentrate by the chlorides that are introduced from the diluate. The final increase of the stack voltage is due to depletion of chloride ions in the boundary layers at the diluate side of the monovalent selective anion exchange membranes. Experiments with model solutions with pure chemicals show a similar behavior.

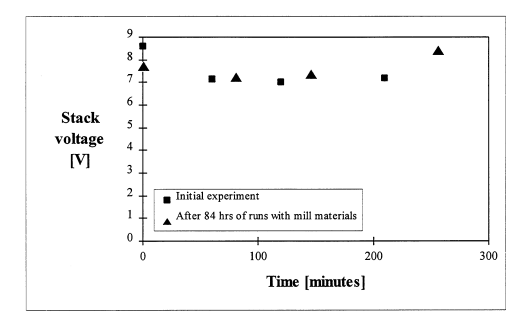


Figure 5: Overall electrical voltage applied at 10 mA/cm². Values for the initial run with mill materials and after a total of 84 hrs of experiments with mill materials.

The result of this extended series of experiments shows that a potentially viable process has been shown here. No pretreatment of any kind was applied to the mill ESP dust solution. Plugging of the flow channels in the system, fouling of the membranes, and precipitation of inorganics could be potential problems. However, no negative impact was observed.

Process economics

A base case bleached kraft pulp mill (BKP mill) with a production of 1000 Air Dry Tons of Pulp per Day (ADTP) was assumed. The energy needed to continuously remove the chloride input to this base case mill by electrodialysis of dissolved ESP dust is estimated below and compared to an energy demand estimate using data from the literature.

1. Assumptions for the base case mill, calculation of energy demand

The source for the total chloride inputs into the liquor system of the base case mill is NCASI (NCASI, 1994). The total chloride input was estimated at 420 kg per day from wood, makeup chemicals, and water. No chloride input from the bleach plant was assumed (no bleach plant effluent recycle to the kraft chemical recovery process). Steady state was assumed (chloride inputs equal chloride purge by electrodialysis). Purge of chloride with the pulp produced was neglected.

Assumptions for the electrodialysis unit were 80% current efficiency for chloride (80% of the electrical current applied transports chloride), 1 m² active cross-sectional membrane area, and 150 concentrate/diluate cell pair units (see Figure 2). The voltage drop per cell pair was assumed to be 2 V (from Figure 5, about 8 V for 4 cell pairs). This is conservative, because the electrode voltage losses in the rather short laboratory stack have a significant impact. This will not be the case in a full-scale unit. The energy for pumping and mixing of the ESP dust with water is not taken into account

Using Faraday's constant F (96,500 coulomb/mol of a univalent salt) and the Chloride input assumed above, an electrical current of 13,214 A results to hold a steady-state chloride concentration in the pulping process. Because multiple use is made of the current passing between the anode and cathode in an electrodialysis stack (150 cell pairs), this value drops to 88 A passing between the anode and cathode. With a current efficiency of 80% for chloride removal, 110 A is the current estimated for the ED unit. This corresponds to 33 kW or 792 kWh/day. With a cost of \$0.026 cents per kWh (assumption for an advantageous utility contract of a pulp mill), the energy cost for the ED unit is \$20.6/day.

2. Comparison with an estimate of the energy demand from published data

The electrical power needed for electrodialytical chloride removal for a 1000 ADTP bleached pulp mill can be roughly estimated to be 751 kWh per day, based on data for similar salt removal levels for water desalination published elsewhere (Baker et al., 1990). The chloride concentration in dissolved ESP dust is quite similar to seawater, and the sulfate and carbonate are assumed here to not influence the separation process.

The result of 751 kWh per day from published data compares very well to the estimate of 792 kWh per day shown above. Because no deterioration in membrane performance was detected in the laboratory experiments, a membrane lifetime of several years should be achieved.

The energy demand for separation would increase essentially proportional with additional chloride input from bleach plant effluent recycle. Membrane area also increases with increased load. In operation, the chloride removal level is easily adjusted through the current passing through the unit. Startup/shutdown of the chloride removal process is very simple.

3. Capital investment

A quotation from a manufacturer of electrodialysis equipment was obtained. The specifications for the base case mill were used. A full-scale electrodialysis unit for the above process was quoted at \$210,000 (electrodialysis unit only, no dissolving equipment or process modifications included). The energy demand for the unit was quoted to be 528 kWh/day by the equipment vendor. This, again, compares well to the estimates above.

Conclusions

Chloride removal from the kraft pulping process under low effluent conditions was the goal of this work. Electrodialysis with monovalent selective membranes was used successfully to separate chloride from a feed solution with a very high concentration of sulfate. The process was shown to operate without significant problems using an actual feed solution containing organic and inorganic contaminants. Electrical current efficiencies above 70% were found. The energy demand of the electrodialysis for a full-scale application was estimated at 1.9 kWh per kilogram of chloride removed.

The work reported here is an example of membrane technology as a kidney for low effluent processing in a major manufacturing industry. The performance of the electrodialysis process with realistic feed solutions is very encouraging. Pilot tests at a kraft pulp mill in the Southeastern United States are underway.

Acknowledgments

The support of this work by the Member Companies of the Institute of Paper Science and Technology is gratefully acknowledged. The donation of membranes by Tokuyama Soda Inc. is also gratefully acknowledged.

Literature Cited

Ahlers, P.-E., Norrstrom, H., Warnqvist, B., "Chlorides in the Kraft Recovery System. Part II: Process and Equipment Aspects on a Closed Bleached Kraft Mill," TAPPI/CPPA Intern. Pulp Bleaching Conf., May 1976, Chicago, p. 19

Baker, R. W., Cussler, E. L., Eykamp, W., Koros, W. J., Riley, R. L., Strathmann, H., "Membrane Separation Systems-A Research and Development Needs Assessment," U.S. DOE report, contract #DE-AC01-88ER30133, vol. II, 1990

Grace, T. M., Leopold, B., Malcolm, E. W. (technical editors), "Volume 5: Alkaline Pulping" in: Kocurek, M. J., Stevens, F., (eds.) "Pulp and Paper Manufacture," Joint Textbook Committee of the Paper Industry of the United States and Canada, Atlanta, 1991

Holman, K. L., Golike, G. P., Carlson, K. R., "Process Simulation of Bleach Plant Effluent Concentrate Recycle to Recovery with Leaching of Chloride from Salt Cake," 1994 Int. Pulp Bleaching Conference, Poster preprints, p. 101

Karjalainen, P. O., Lofkrantz, J. E., Christie, R. D., "Chloride Buildup in Kraft Liquor Systems," Pulp & Paper Canada, 73(12):95, 1972

Keitaanniemi, O., Virkola, N.-E., "Undesirable Elements in Causticizing Systems," TAPPI Journal, 65(7):89, 1982

Maruko, S., Iwadare, T., Kobuchi, Y., Motomura, K., Matsunaga, Y., "Method of Dechlorination," Jap. Patent No. 22,051-1980

Moy, W. A., Joyce, P., Styan, G. E., "Removal of Sodium Chloride from Kraft Recovery Systems," Pulp & Paper Canada, 75(4):88, 1974

NCASI (National Council of the Paper Industry for Air and Stream Improvement, Inc.), "A Study of Kraft Recovery Furnace Hydrochloric Acid Emissions," NCASI Technical Bulletin No. 674, 1994

Rapson, W. H., "The Feasibility of Recovery of Bleach Plant Effluent to Eliminate Water Pollution by Kraft Pulp Mills," Pulp & Paper Canada, p. T635, Dec. 1967

Reeve, D. W., "The Effluent-free Bleached Kraft Pulp Mill. Part VII: Sodium Chloride in Alkaline Pulping and Chemical Removal," Pulp & Paper Canada, 77(8):35, 1976

Reeve, D. W., Tran, H. N., Barham, D., "The Effluent-free Bleached Kraft Pulp Mill - Part XI," Pulp & Paper Canada, 82(9):105, 1981

Saracco, G., Zanetti, M. C., Onofrio, M., "Novel Application of Monovalent-Ion-Permselective Membranes to the Recovery Treatment of an Industrial Wastewater by Electrodialysis," Ind. Eng. Chem. Res. 1993, 32, 657-662

Tran, H. N., "How does a Recovery Boiler Become Plugged?", TAPPI Journal, 69(11):102, 1986

Wright, R. H., "The Separation of Sodium Chloride from Precipitator Salt-Cake," Pulp and Paper Magazine of Canada, 57(12):171, Nov. 1956

·			

		•		
-				