BLEACHING CHEMICALS FOR THE KRAFT PULPING INDUSTRY

The pulp used to make paper is brown when first produced. For most applications this is unacceptable, so it is bleached to make it white. The chemicals used to do this are chlorine (Cl₂), caustic soda (NaOH) and chlorine dioxide (ClO₂).

Chlorine and caustic soda are produced by electrolysis of solutions of NaCl. In this solution it is the water, not the Na⁺ ion, that is reduced. The two solutions are separated by a membrane that allows Na⁺ ions but not OH⁻ ions to pass through, so that the chlorine and caustic soda do not mix. The reactions occurring are:

At the anode: \[ \text{2Cl}^- (\text{aq}) \rightarrow \text{Cl}_2 (\text{g}) + 2\text{e}^- \]

At the cathode: \[ \text{2Na}^+ + 2\text{H}_2\text{O} + 2\text{e}^- \rightarrow 2\text{NaOH} + \text{H}_2(\text{g}) \]

Overall: \[ 2\text{NaCl} + 2\text{H}_2\text{O} + \text{electrical energy} \rightarrow 2\text{NaOH} + \text{Cl}_2(\text{g}) + \text{H}_2(\text{g}) \]

The chlorine dioxide is produced in a three step process using the reactions given below.

Step 1 \[ \text{NaCl} + 3\text{H}_2\text{O} + \text{electrical energy} \rightarrow \text{NaClO}_3 + 3\text{H}_2 \]

Step 2 \[ \text{H}_2 + \text{Cl}_2 \rightarrow 2\text{HCl} \]

Step 3 \[ \text{NaClO}_3 + 2\text{HCl} \rightarrow \text{ClO}_2 + \frac{1}{2}\text{Cl}_2 + \text{NaCl} + \text{H}_2\text{O} \]

Overall \[ \frac{1}{2}\text{Cl}_2 + 2\text{H}_2\text{O} \rightarrow \text{ClO}_2 + 2\text{H}_2 \]

In addition to producing aqueous Cl₂, NaOH and ClO₂, the plant produces excess NaClO₃ and HCl, and produces NaOCl and liquid Cl₂ as byproducts. These are sold for making commercial bleaches and for water treatment.

INTRODUCTION

While unbleached kraft pulps are strong, their brown colour limits their use to grades which do not require a white fibre. Kraft pulps can be bleached to produce a white fibre. Bleaching kraft pulps can be done in a number of ways, in New Zealand bleaching is done using a combination of chlorine dioxide (ClO₂), oxygen (O₂) and caustic soda (NaOH, also known as sodium hydroxide).

Caustic soda is produced by electrolysis of brine (aqueous sodium chloride), and the co-produced chlorine can be used to manufacture the chlorine dioxide required by the bleaching plant. Oxygen can be produced either cryogenically (cooling air until it liquifies and using the boiling point difference to separate O₂ from N₂) or using the PSA process (see article).

The processes involved in making Cl₂, NaOH and ClO₂ also lead to the production of HCl, NaOCl and NaClO₃ as well. Some of these are used on site, and some are sold.
Uses of products and byproducts

Chlorine, Cl₂
Chlorine is used in liquid form for pulp bleaching. It is also a starting material for HCl and NaOCl production. Liquid chlorine is sold for uses such as water treatment.

Sodium hydroxide, NaOH
All the caustic soda produced is used for pulp bleaching.

Chlorine dioxide, ClO₂
All the chlorine dioxide produced is used for pulp bleaching.

Hydrochloric acid, HCl
Internally, hydrochloric acid is used in ClO₂ manufacture, pH control and the regeneration of ion exchange resins (see article). It is also sold to the metals industry (for metal cleaning and pickling) and the chemical cleaning industry.

Sodium hypochlorite, NaOCl
This is one of the active ingredients of commercial bleaches such as "Janola". Excess chlorine is used to make NaOCl for sale to bleach manufacturers.

Sodium chlorate, NaClO₃
This is a starting material for ClO₂ manufacture. Any excess is sold to other manufacturers for a variety of uses including the production of weed killer, matches and fireworks and the refining of uranium ore.

THE CHLORINE/CAUSTIC SODA MANUFACTURING PROCESS

These two chemicals are produced concurrently. The three processes outlined below have all been used to produce them commercially, although only the latter process is still in use in New Zealand.

The mercury cell process
This is the oldest process. In this process the electrolyser cell consists of an anode (made of graphite or titanium) and a cathode formed by mercury. Brine is introduced into the cell. Cl₂ gas forms at the anode and bubbles out of solution. At the cathode, sodium (Na) forms an amalgam with the mercury cathode. The sodium/mercury amalgam is removed from the cell and is reacted with water to form caustic soda (NaOH) and hydrogen gas (H₂). The main disadvantage of this process is that the products and plant effluent are contaminated with mercury, and for this reason all mercury cell electrolysers were decommissioned many years ago.

The diaphragm process
A brine solution is pumped into the cell where the anode and cathode are separated by an asbestos diaphragm. Cl₂ gas is formed at the anode, and the NaOH is recovered from the solution leaving the cell (a mixture of depleted brine and NaOH). The main disadvantage of this process is that the NaOH has to be recovered by evaporation which requires a further large energy input.
The membrane cell process
This process was developed to eliminate the use of mercury and to reduce the energy required to produce the NaOH solution. It was developed by Asahi Chemical Industry Co Limited of Japan. Here a titanium anode and a steel cathode are used, separated by a cation exchange membrane which allows Na⁺ ions to migrate to the cathode, but prevents OH⁻ ions from going the other way. This results in the formation of Cl₂ at the anode, and NaOH on the cathode side.

Step 1 - Anolyte preparation
The solution that enters the electrolysis cell on the anode side is called the "anolyte". It is essentially a very pure solution of brine (water saturated with NaCl). Anolyte is prepared as follows.

Depleted brine from the electrolysis cells is kept in a storage tank. This is a solution of NaCl dissolved in water, but it is below the saturation point. The first stage of anolyte preparation is to remove some of this solution from the anolyte tank and pump it through a bed of raw salt in the saturator. This has brine added to the bottom and salt added to a hopper from the top.

This brine contains small amounts of metal cations such as Ca²⁺, Ni²⁺ and Fe³⁺, as well as the Na⁺. These must be removed before the solution goes to the electrolysis cell to prevent the formation of insoluble hydroxides which would clog up the electrolyser membrane. They are removed by adding various chemicals including barium carbonate (BaCO₃), sodium carbonate (Na₂CO₃) and ferric chloride (FeCl₃), which cause the cations to precipitate out. The precipitates are then removed in a settling tank. The brine is then filtered to remove colloidal precipitates and passed through two cation exchange columns. Metal ions (except Na⁺) are removed to form a very pure solution of NaCl. Finally a small amount of HCl is added, to neutralise any OH⁻ that gets into the solution during electrolysis.

Step 2 - Catholyte production
Catholyte is a solution of NaOH (about 19.4 wt% NaOH). It is prepared by pumping back most of the NaOH-containing solution from the electrolyser, and diluting it back to 19.4% NaOH with water. The water used is deionised water that has been used to scrub the hydrogen gas formed in the electrolysers.

Step 3 - Electrolysis
The anolyte and catholyte solutions then enter the electrolysis cell. The anolyte enters on the anode side of the membrane at the bottom corner of the cell and exits from the opposite top corner of the cell. Similarly, the catholyte also enters at the bottom of the cell, but on the cathode side. Inside the cell chloride ions are oxidised to form chlorine gas and the water on the cathode side is reduced to form hydroxide ions, as given in the reactions below:

At anode: \[ 2\text{Cl}^- (aq) \rightarrow \text{Cl}_2(g) + 2e^- \]
At cathode: \[ 2\text{Na}^+ + 2\text{H}_2\text{O} + 2e^- \rightarrow 2\text{NaOH} + \text{H}_2(g) \]
Overall: \[ 2\text{NaCl} + 2\text{H}_2\text{O} + \text{electrical energy} \rightarrow 2\text{NaOH} + \text{Cl}_2(g) + \text{H}_2(g) \]

The electrolysis cell
The electrolysis takes place in electrolysers consisting of 42 cells in series. The cells are held in a frame, with one side of the cell frame being the anode (made of titanium) and the other
being the cathode (made of carbon). A large DC current flows from the cathode to the anode along the length of the electrolyser.

As electrolysis takes place the sodium ions are attracted to the cathode and the OH\(^{-}\) ions are attracted to the anode. However, to ensure that NaOH and Cl\(_2\) do not react together to form sodium hypochlorite, the two chemicals are produced in compartments separated by a cation exchange membrane that allows Na\(^{+}\) ions through, but restricts OH\(^{-}\) migration. This means that Cl\(_2\) forms on the anolyte side and NaOH forms on the catholyte side. A small amount of OH\(^{-}\) does pass through the membrane, but this is neutralised by the acid that was added to the anolyte. The membrane used at Carter Holt Harvey Pulp & Paper’s Kinleith mill is a perfluoro carboxylic acid membrane with the general chemical structure given below.

\[
\begin{align*}
\text{---CF}_2\text{--CF}_2\text{--CF--CF}_2\text{---} \\
\text{O} \\
\text{CF}_2 \\
\text{CF}---\text{CF}_3/n \\
\text{O} \\
\text{CF}_2 \\
\text{COOH}
\end{align*}
\]

**Figure 1** gives a schematic representation of an electrolysis cell. Note that \(\eta\) refers to current efficiency, the fraction of the current that is actually used to produce Cl\(_2\). This is less than one because the oxidation of water to oxygen competes with the chlorine oxidation at the anode.

Some details of the electrolyser operating conditions are given below:

- Number of Electrolysers: 2
- Number of Cells per Electrolyser: 42
- Power Supply: 20 kA DC at 160 V. The electrolysers are operated in parallel and receive 10 kA each.
- Cell voltage: 3.0 - 4.0 V
- Power consumption: 2200 - 2400 DC kWhr/t NaOH

**Step 4 - Cl\(_2\) and NaOH recovery**

The anolyte leaving the cell is a two phase mixture of depleted brine and chlorine gas. A cyclonic separator is used to separate the chlorine gas from the solution and the depleted brine is returned to the anolyte tank. The chlorine gas is scrubbed by incoming anolyte solution, and then used to form liquid chlorine, sodium hypochlorite and hydrochloric acid.

A cyclonic separator separates the caustic solution from the hydrogen gas. The caustic solution returns to the catholyte tank and the hydrogen gas enters the bottom of the scrubber to be scrubbed by incoming catholyte solution. The caustic is now ready to be used for paper bleaching, and the hydrogen gas is used later on in the process for HCl production.
Chlorine/Caustic balance
As shown above the overall reaction to produce Cl₂ and NaOH is:

\[
2\text{NaCl} + 2\text{H}_2\text{O} + \text{electrical energy} \rightarrow \text{Cl}_2 + 2\text{NaOH} + \text{H}_2
\]

Now the species involved have the following molecular weights:
- NaCl = 58.5 g mol⁻¹
- H₂O = 18 g mol⁻¹
- Cl₂ = 71 g mol⁻¹
- NaOH = 40 g mol⁻¹
- H₂ = 2 g mol⁻¹

Thus for every tonne of Cl₂ produced, 1.13 tonnes of NaOH is also produced. This means that the pulp mill must either keep the NaOH:Cl₂ consumption ratio very close to 1.1:1, or purchase extra NaOH. Overseas, pulp mills generally purchase all the NaOH and Cl₂ that they need, but New Zealand is a sufficiently far away from commercial chlor/alkali plants to make importing these chemicals prohibitively expensive. Hence the amount of NaOH able to be produced is strictly limited by the amount of Cl₂ that can be used for bleaching, or for producing other chemicals, or which can be sold. If the 1.1:1 ratio is exceeded, expensive imported caustic soda must be purchased.
THE CHLORINE DIOXIDE MANUFACTURING PROCESS

Chlorine Dioxide (ClO₂) has been used as a bleaching chemical commercially for many years. The first ClO₂ plants were installed at overseas mills in 1946. Over the years a large number of processes have been developed to produce ClO₂, and these are summarised in Table 1. All of the processes are based on the reduction of chlorate (using Cl⁻, CH₃OH or SO₂) in the presence of either sulphuric or hydrochloric acid.

### Table 1 - ClO₂ Production Processes

<table>
<thead>
<tr>
<th>Process</th>
<th>Reducing agent</th>
<th>Acidity</th>
<th>Byproducts</th>
<th>Yield / %</th>
</tr>
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<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Cl₂¹</td>
<td>Na₂SO₄</td>
</tr>
<tr>
<td>R2</td>
<td>NaCl</td>
<td>High</td>
<td>0.6</td>
<td>6.9</td>
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<td>R2H</td>
<td>HCl/NaCl</td>
<td>High</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Mathieson</td>
<td>SO₂</td>
<td>High</td>
<td>—</td>
<td>3.4</td>
</tr>
<tr>
<td>Salted</td>
<td>SO₂/NaCl</td>
<td>High</td>
<td>0.15</td>
<td>4.7</td>
</tr>
<tr>
<td>Matheson</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Solvay</td>
<td>CH₃OH</td>
<td>9 - 11N</td>
<td>—</td>
<td>3.5</td>
</tr>
<tr>
<td>Salted</td>
<td>CH₃OH/HCl</td>
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<td>0.12</td>
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<tr>
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<td></td>
<td></td>
</tr>
<tr>
<td>R8</td>
<td>CH₃OH</td>
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<td>—</td>
<td>1.5</td>
</tr>
<tr>
<td>R3/SVP</td>
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<td>2.3</td>
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<tr>
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<td>Na₂SO₄</td>
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<td>1.4</td>
</tr>
<tr>
<td>Integrated</td>
<td>HCl/NaCl</td>
<td>Low</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

Notes:
⁺ Produces acid effluent
⁺⁺ Has no byproducts

The early processes (eg R2, Mathieson, Solvay) produce a highly acidic effluent (which can be troublesome to dispose of), together with large amounts of saltcake (Na₂SO₄). The acid effluent can be added to black liquor (see section on kraft pulping) as a make up chemical in the regeneration stage provided that the amount of Na₂SO₄ does not exceed the mills requirements and that precautions are taken to minimise emissions of H₂S when the acid reacts with sulphides in the black liquor. However, these conditions are quite hard to meet and these processes are now out of favour because of the high production of Na₂SO₄.

¹ Cl₂ and Na₂SO₄ quantities given in tonnes of Cl₂ or Na₂SO₄ per tonne of ClO₂ produced.
At Kinleith two processes have been used. From 1972 to 1991 the SVP process (also known as the R3 process) was used. The basic reaction occurring here is:

\[ \text{NaClO}_3 + \text{NaCl} + \text{H}_2\text{SO}_4 \rightarrow \text{ClO}_2 + \frac{1}{2}\text{Cl}_2 + \text{Na}_2\text{SO}_4 + \text{H}_2\text{O} \]

There is a competing reaction which produces no \( \text{ClO}_2 \):

\[ \text{NaClO}_3 + 5\text{NaCl} + 3\text{H}_2\text{SO}_4 \rightarrow 3\text{Cl}_2 + 3\text{Na}_2\text{SO}_4 + 3\text{H}_2\text{O} \]

Catalysts are added to encourage the \( \text{ClO}_2 \) producing reaction.

From 1991 the process used in New Zealand has been the integrated process, which involves producing sodium chlorate and HCl and then reacting them together to form ClO2. This process is not used much internationally because of the high capital costs of the electrolyser, but interest in the process is increasing. It is ideal for New Zealand because it allows the Kinleith mill to be completely self sufficient for NaOH while providing the bleaching chemicals necessary to bleach pulp. The reactions involved are given below:

\[ \text{NaCl} + 3\text{H}_2\text{O} + \text{electrical energy} \rightarrow \text{NaClO}_3 + 3\text{H}_2 \]

\[ \text{H}_2 + \text{Cl}_2 \rightarrow 2\text{HCl} \]

\[ \text{NaClO}_3 + 2\text{HCl} \rightarrow \text{ClO}_2 + \frac{1}{2}\text{Cl}_2 + \text{NaCl} + \text{H}_2\text{O} \]

\[ \text{Cl}_2 + \text{H}_2\text{O} \rightarrow \text{ClO}_2 + 2\text{H}_2 \]

**Step 1 - Sodium chlorate production**

Sodium chlorate is produced by electrolysing salt using a chlor/alkali cell without the cation exchange membrane. The overall reaction occurring is:

\[ \text{NaCl} + 3\text{H}_2\text{O} + \text{electrical energy} \rightarrow \text{NaClO}_3 + 3\text{H}_2 \]

Some of the operating conditions for the integrated plant are listed below:

- Chlorate Cell Power Supply 57 kA DC at 135 V
- No of cells 40
- Cell Voltage 2.0 - 3.0 V
- Power Consumption about 5000 kWhr/t chlorate

**Step 2 - HCl production**

Hydrogen chloride is formed by burning chlorine in a slight excess of hydrogen without oxygen. The gases are originally obtained in the chlor/alkali cells, and are regenerated at various subsequent process steps such as chlorate production.

\[ \text{H}_2(g) + \text{Cl}_2(g) \rightarrow 2\text{HCl}(g) \]

The hydrogen chloride gas is absorbed in demineralised water to form 33wt% hydrochloric acid.

**Step 3 - Chlorine dioxide generation**

Chlorate solution and 33 wt% HCl are mixed in the generator to produce ClO2 and Cl2. The ClO2 is extracted by preferential absorption into chilled water. Some Cl2 is recycled to the generator where together with air it is used to dilute the ClO2 to reduce the concentration of
ClO₂ and hence the likelihood of decomposition. The remaining Cl₂ is added to the Cl₂ from the chlor/alkali plant to make the required HCl. As the reaction in the generator proceeds NaCl builds up in solution. This is sent back to the chlorate cells where it is converted back by electrolysis to NaClO₃.

PLANT SAFETY

Chlorine dioxide
Chlorine dioxide is a very unstable compound, it is considered too hazardous to transport. For this reason chlorine dioxide is always produced on the site where it is used. The parameters which must be controlled to minimise decomposition of the ClO₂ are:

i) keep concentration low (typically < 10% by volume)
ii) shield from ultraviolet (uv) light
iii) keep uncontaminated
iv) keep cool

Sodium chlorate
Cryatallised chlorate has to be handled with care. When heated dry chlorate releases O₂, which can lead to spontaneous combustion and explosions. Once alight, a fire accelerated with chlorate cannot be put out by smoothering (eg CO₂ extinguisher), but must be quenched with large amounts of water.

ANCILLIARY PROCESSES

Liquid chlorine production
The Cl₂ gas from the anolyte tank is saturated with water, making the Cl₂ very corrosive to steel. As the equipment used to process and store liquid chlorine is made from steel the first step in processing the Cl₂ for liquefaction is to remove the water from the gas. The water is removed by contacting the gas with concentrated sulfuric acid (about 98 wt% H₂SO₄) which absorbs the acid. The Cl₂ is then compressed to 550 kPa using a two stage reciprocating compressor to reduce the temperature at which it will liquefy. The gas is then cooled in a heat exchanger (using an ammonia refrigeration system) to produce liquid chlorine. As ammonia reacts violently with chlorine, the ammonia is used to cool kerosene which is in turn used to cool the Cl₂ to produce liquid chlorine. In this way the chance of an explosion if the heat exchanger leaked is minimised.

Sodium hypochlorite production
Cl₂ not used to make HCl and liquid chlorine is made into sodium hypochlorite (hypo²) solution. The Cl₂ gas is reacted with NaOH to form the hypo in a very fast, very exothermic reaction as follows:

\[
Cl_2 + 2NaOH(aq) \rightarrow NaOCl(aq) + H_2O
\]

The hypo solution is produced at a concentration about 165 g available Cl₂ L⁻¹.

²Note that in this context "hypo" refers to sodium hypochlorite while in photography it refers to sodium thiosulfate.
The hypo system is used to remove chlorine from process streams before they are vented to atmosphere and is also used to safely dispose of chlorine during process upsets such as shut downs and power failures.

**THE ROLE OF THE LABORATORY**

The chemical plant is an area where chemical reactions and electrochemistry are applied in continuous processes. Regular monitoring of these reactions is vital, and therefore the plant has it’s own laboratory, and testing is carried out by plant operators around the clock. Many of the processes in each plant are similar with routine testing including brine strength, control of brine treatment systems, safety gas checks, strength of caustic soda, hydrochloric acid, sodium hypochlorite, sulphuric acid in chlorine driers, scrubbers, filters, ion exchanger regeneration chemicals and so forth.

The Kinleith Mill has a central Analytical Laboratory capable of carrying out a wide range of testing for all areas of the site it services. This laboratory interfaces with the chemical plant by providing test results which would be outside the scope of a plant lab to perform, or which uses instrumentation unsuitable for an industrial environment.

**Chlor-alkali plant**

Routine analyses include ppb level determination of metal ions in the purified brine. Elements such as Ca, Mg, Ba, Fe and Ni can “poison” the ion exchange membranes in the electrolysers, and these are monitored by daily checks on the brine feed and anolyte circulation. Currently graphite furnace atomic absorption spectrophotometry is used. The high sodium chloride concentrations (150 - 300 gram per litre) and low detection limit requirements (1 ppb) call for careful analysis programs to minimise interference effects. Elevated levels (above 20ppb) of these metals are indicative of a problem with the ion exchange columns, or contamination from additives or corrosion. Hydrochloric acid and demineralised water purity are monitored as these enter the brine circuit also.

By-products from the chemical plant are sold to chemical suppliers, local authorities and other manufacturers. The laboratory tests these chemicals as they are despatched to check conformity to specification, and reports results to customers.

**Chlorate & Chlorine Dioxide Plant**

The chlorate and chlorine dioxide processes are more tolerant of contamination, but are less tolerant of changes in the process conditions. The plant carries out most of it’s own testing to ensure the conditions are maintained.

During commissioning and plant optimisation the laboratory profiles the chlorine dioxide generator, analysing inputs, and compartment by compartment composition. This includes chlorate, chloride, acid strength, dissolved gases, and gas composition. From this information decisions are made on temperature, dilution air, and feed ratios. Mass balances and overall plant efficiency can be derived. Wet chemistry methods are used for these analyses. Sampling technique is critical, all samples are diluted directly into chilled water to minimise gas-off and to quench the reactions in order to “freeze” the chemical conditions in the compartment at the time.
The Analytical Laboratory checks raw materials and water supplies, and trouble shoots problems in both plants. Chemists work with the plant staff to optimise the process, reduce losses, and improve reliability of equipment. Test chemicals and standard solutions are supplied to the plant lab from here also.

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