The Electro-oxidation of Lignin in Sappi Saiccor Dissolving Pulp Mill Effluent

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Abstract

Electro-oxidation reactions using a nickel anode were carried out on the calcium-spent liquor effluent obtained from Sappi Saiccor (formerly South African Industrial Cellulose Corporation) dissolving pulp mill as well as on lignin and lignan type compounds previously identified in the effluent. Voltammograms were obtained for each solution in order to identify the oxidation potentials of the compounds to be electro-oxidized. Value-added products such as vanillin and syringaldehyde were identified in the electro-oxidized reaction mixtures using gas chromatography-mass spectrometry (GC-MS) and high performance liquid chromatography (HPLC). Profiles of the changes in concentration of these compounds were determined as a function of time with maximum concentrations reached within the first half hour. These findings are significant in that few electro-oxidation reactions have been carried out on the effluent of a pulp mill which uses the acid bisulfite pulping process and no results have previously been reported on the electro-oxidation of syringaldehyde. This study contributes to a further understanding of the electro-oxidation of lignin and is of value to the paper and pulp industry at large. Cleaning of acidic pulp effluent by electro-oxidation is shown to be effective.

Keywords: electro-oxidation, lignin, effluent, vanillin, syringaldehyde

1. Introduction

Sappi Saiccor, situated near Durban, South Africa, is one of the few pulp mills that still partially use calcium bisulfite as the base for the acid bisulfite pulping process. This process produces a high-grade cellulose pulp but significant volumes of effluent waste are also generated, which are discharged to sea via a 7 km pipeline. Environmental regulations require that effluent reductions are made and therefore Saiccor has embarked on an investigation of the effluent itself.

Research on the effluent has shown it to contain the lignan isomer syringaresinol, as the major constituent, some lignin-type precursors, fatty acids, sugars and lignosulfonates (Moodley et al., 2003). The aim of this project was to oxidize the effluent to produce value-added products that could be extracted for commercial purposes, thereby reducing the amount of effluent pumped out to sea as waste

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as well as investigating the electrochemical cleaning of the effluent. This is the first attempt at performing any type of oxidation on Sappi Saiccor’s effluent and the results could be of value to the industrial paper and pulp community. Previous work on the oxidation of lignin was based primarily on Kraft pulping effluent, which is alkaline, compared to our acid bisulfite pulping effluent. Furthermore, Saiccor uses hardwoods with a higher syringyl content compared to mills that use softwoods which have higher guaiacol content. Thus the source of the wood and the pulping process used combine to produce an effluent unique to Saiccor. Previous electro-oxidation reactions have been done on pure compounds but not on a mixture of compounds obtained from effluent as carried out in this work. The results presented in this paper for the electro-oxidation of syringaldehyde are also the first reported results in literature.

Many chemical oxidants have been employed for the oxidation of lignin. Although nitrobenzene is a suitable oxidizing agent, high temperatures and pressures are necessary and the formation of nitro derivatives make these oxidations commercially and environmentally undesirable. The use of oxygen as an oxidant also requires high temperatures and pressures which again make it an undesirable process. Reactions with hydrogen peroxide tend to form acids and often lead to degradation of the aromatic ring. Thus electrolysis of lignin could be an attractive alternative in the oxidation of lignin and lignosulfonates as it offers several advantages over the traditional chemical oxidation processes. In particular, electrolysis of lignin and lignosulfonates in basic media can use relatively lower temperatures and pressures compared to reactions with oxygen and nitrobenzene and there is better control of the reaction, leading to greater selectivity (Varey, 1987; Utley, 1986, Utley, 1987, Utley, 1988). Degradation of lignin often produces products such as aldehydes, ketones and acids.

Some of the earliest studies on the electro-oxidation of organic molecules involved phenolics and it was found that electro-oxidation was comparable to chemical oxidation reactions, forming the aromatic aldehyde and acid (James, 1899; Law, 1906). Electro-oxidation of phenolics and lignin model compounds were investigated with the aim of using the technique as an identification tool and for the oxidation and synthesis of phenolics and lignin derivatives such as vanillin and syringic acid (Hedenberg, 1953; Gaylor et al., 1953; Lund, 1957; Vermillion, 1963; Kovalenko et al., 1973; Kovalenko et al., 1976; Kovalenko et al., 2000; Demin et al., 1980). Various graphite and platinum electrodes were investigated and it was realized fairly early that the nature of the electrode surface played an important role (Hedenberg, 1953; Gaylor et al., 1953; Gaylor et al., 1957; Elving, 1958; Morris, 1959). Many of these studies observed the formation of a thin surface film which was thought to be the result of polymerization and recent studies show a possible application of it in the protection of steel (Vagin et al., 2006). Electro-oxidation of lignin type compounds in base using platinum and nickel electrodes showed the formation of dimers using starting materials obtained from an acidic media (Pearl, 1952), which has also been noted in the present study.

Smith and Utley showed that a combination of nitroaromatics in an electrolysis reaction of lignin in base required lower temperatures than either reaction on its own (Smith, 1989). The disadvantage was
that nitro derivatives were still present in the end product. A patent by Utley and Smith claimed to produce yields of useful monomers, greater than 10%, by passing an electric current through an alkaline solution of lignin at temperatures above 100°C with continuous mixing (Utley, 1988). This method claimed to produce vanillin or partial cleavage products such as dimers, trimers or oligomers of monocyclic species. Studies of the mechanism of electro-oxidative cleavage of dimers have shown that aldehydes such as vanillin and syringaldehyde form as well as carboxylic acids (Pardini et al., 1991). These products all formed at a nickel anode resulting from an oxygenated nickel (III) anode surface (Pardini et al., 1991). Recent research on the combination of photochemical and electrochemical oxidation methods on Kraft lignin showed the production of vanillin and vanillic acid (Tian et al., 2010).

Studies were carried out on the influence of different electrode materials such as Pt, Au, Ni, Cu, DSA-O₂ (dimensionally stable anode), PbO₂ and boron-doped diamond electrodes, on phenolics and lignin electro-oxidation reactions (Kovalenko et al., 1977; Parpot et al., 2000; Canizares et al., 2002; Iniesta et al., 2001). Early studies on graphite, copper, platinum and nickel anodes showed that platinum and nickel produced higher amounts of aldehyde and a comparison of the nickel and DSA-O₂ anodes at the same high current densities of 750 A/m², showed that nickel produced a higher concentration of vanillin per unit of charge used (Kovalenko et al., 1977; Parpot et al., 2000).

Current research on the electrochemical oxidation of Kraft lignin has involved the use of a number of different natural and synthetic compounds that may act as redox mediators or enhancers during these electrochemical oxidation reactions of the Kraft lignin (Arzola et al., 2009). The use of modified N-hydroxyphthalimide has also found to be beneficial in promoting electrochemical oxidation of non-phenolic lignin model compounds in the presence of a laccase mediator (Annunziatini et al., 2005).

2. Experimental

2.1 Pre-treatment procedure

The nickel anode, which forms a nickel (III) peroxide surface, was cleaned by holding the electrode at negative, and thereafter positive, potentials for 30 seconds each, rinsed with Millipore water and then placed in an ultrasonic bath for 5 minutes, followed by cleaning in nitric/sulfuric acid mixtures. A nickel working electrode, platinum counter electrode and a Ag/AgCl reference electrode were connected to a BAS potentiostat CV-27. The following figures show the schematic diagram and the actual electro-oxidation setup.
2.2 Solution preparation and voltammetry

Standard solutions containing pure syringaldehyde (0.01 M) in KOH (0.1 M) and sodium perchlorate (0.5 M) were prepared in Millipore water. Background voltammograms were obtained for 0.5 g/ℓ solutions of pure compounds in KOH and sodium perchlorate (pH 12.45) with continuous stirring. The solutions were electro-oxidized at the suitable oxidation potentials for a period of time with continuous stirring. The oxidized solution was extracted with chloroform in a sulfuric acid medium. All chemicals used were analytical grade.

2.3 Analysis

The end product was analyzed using GC-MS and HPLC. GC-MS analysis involved 1μℓ manual injections onto a HP5-MS column with 1:75 split. The sample was held for two minutes at 50°C, the
temperature was ramped at 20°C per minute until it reached 300°C and thereafter the sample was held for 20 minutes. Low-resolution mass spectrometry was performed on an Agilent 5973 mass spectrometer connected to a 6890GC. The results of the mass spectrum were matched with mass spectra of NIST reference standards.

HPLC analysis was performed on an Agilent system with an UV detector. Filtered samples (25 μℓ in acetonitrile) were introduced onto a ZORBAX C-18 extend column connected to a C-8 column in the LC. The solvent system was increased from 0-100% acetonitrile with 0.2% formic acid in 60 minutes with a flowrate of 1.000 mL/min and the oven temperature was maintained at 40°C. Some analyses were conducted using a gradient system of 0-100% acetonitrile with 0.2% formic acid in 170 minutes with the same flowrate and oven temperature as above but with a stop-time of 60 minutes and a 10 minute flush between samples. Blanks were obtained according to standard procedures. Quantitative analyses were performed using calibration standards with the inclusion of a 0.002 M internal standard, 3, 4–dimethoxybenzaldehyde in a solution of KOH (0.1 M) and NaClO₄·H₂O (0.5 M), in each standard. For the electro-oxidation reactions, samples (1.5 mL) were taken every hour during the electro-oxidation of syringaldehyde and every half hour for the remaining reactions, filtered and placed in vials containing 30 μℓ of filtered 0.1 M internal standard. The samples were analyzed up to 5 times using the LC instrument and concentrations were calculated from the calibration curves. These values were used to calculate the mass of compound produced per gram of starting material. The initial mass of the compound already present per gram of starting material was determined at time = 0 and was subtracted from the mass of compound produced per gram of starting material calculated at each sample time. The resulting mass of compound produced per gram of starting material was converted to a percentage and the average determined. This result is referred to as percentage concentration and will be discussed later.

3. Results and discussion

Electro-oxidations were performed on standard solutions of syringaldehyde and the lignan syringaresinol which are known to be present in the effluent, as well as on the organic extracted portion of the effluent, a concentrated form of the effluent and lignosulfonates obtained from Lignotech, all originating from an acid bisulfite pulping process. Cyclic voltammograms (CV) or linear scans were obtained for each solution including the background electrolyte on its own. The linear scans (Figure 3) for 0.5 g/L of syringaldehyde solution (dashed line) showed an oxidation peak potential at 0.66 V vs Ag/AgCl compared to the oxidation peak for the background electrolyte on its own (solid line).
After electro-oxidation for 5 hours at 0.66 V, analysis of the end product showed the presence of syringic acid together with other dimers and compounds (Figure 4). The presence of syringic acid in the final mixture confirmed that electro-oxidation had occurred. The identification of each peak in the GC spectrum was based on its mass spectrum and the structures of the dimers were confirmed by comparison of the mass spectra with published data (Majumder, 1993; Bangani, 1998; Pearl, 1957).
Figure 5: Structures of compounds identified after electro-oxidation of syringaldehyde

The phthalates identified are possibly from the plastic fittings used during the reaction. A quantitative analysis of the reaction product showed that the maximum conversion of syringaldehyde to syringic acid occurred within the first 2 hours of electro-oxidation yielding 0.28% (of starting material) ($\pm 0.48 \times 10^{-2}$%) of syringic acid (Figure 6). Thereafter, the percentage concentration of syringic acid decreased suggesting a method of effluent cleaning. In this work, the percentage concentration is the mass percentage concentration of the compound formed from the starting material. It is calculated from the following formula:

$$\frac{M_t - M_i}{M_s}$$

Where, $M_t$ is the mass of the compound at time $t$, $M_i$ is the mass of the compound at time 0 and $M_s$ is the mass of the starting material.

Since the definition of percentage concentration used in this work is based on the additional mass of compound produced per gram of starting material, then if the additional mass produced decreases over time due to electro-degradation, the percentage concentration will decrease. Furthermore, the initial starting material (in the case of the concentrated effluent, organic portion of the effluent and the lignosulfonate mixture) already contains a certain mass of the compound of interest which may be destroyed or more may be produced during the reaction. This initial starting material must be
subtracted from the mass of compound produced at each sample time to give a true reflection of the actual amount of compound produced from the reaction itself. If the overall mass of compound per gram of starting material present at each sample point decreases and the initial mass of compound present is subtracted, negative concentrations may be obtained because the change in mass of compound produced per gram of starting material is plotted against time, and not the total mass of compound produced per gram of starting material. These reduced values will also be referred to as percentage concentrations in order to maintain consistency and allow for comparison of different compounds.

Figure 6: Formation of syringic acid during electro–oxidation of syringaldehyde

Studies on the effluent identified the lignan syringaresinol as the major component and therefore electro-oxidation reactions were carried out on this compound (Moodley et al., 2003). A linear sweep of a 0.5 g/l solution of the lignan showed oxidation peaks at 0.35 V vs Ag/AgCl and 0.59 V vs Ag/AgCl (Figure 7). This suggested that the oxidation of syringaresinol proceeds via a two-step process that forms an intermediate, which is then oxidized to the final product (Pearl, 1957; Yoshida, 1984).

Figure 7: Oxidation potentials for the lignin, syringaresinol at pH 12.45
Qualitative analysis of the electro-oxidized extract of the lignan by GC-MS (Figure 8) showed the presence of a lignan derivative, some phthalate contaminants, a small amount of syringaldehyde and an even smaller amount of syringic acid whose peak is not clearly seen in Figure 8. The structures of compound A and the lignan derivative were not conclusively identified as insufficient quantities of these compounds were obtained.

![Figure 8: GC spectrum of the electro-oxidized lignan](image)

The quantitative analysis of syringaldehyde and syringic acid formed during the reaction showed that the concentration of syringaldehyde increased slightly and thereafter stabilized after about 2.5 hours (Figure 9). Syringic acid concentration on the other hand decreased. In this case a small amount of syringic acid forms during the pre-treatment† stage and is subtracted from syringic acid formed during the reaction. Since the total amount of syringic acid also decreased as the reaction proceeded and together with the very small amount formed initially during the reaction, the eventual change in mass of syringic acid produced per gram of starting material is a “negative or reduced percentage concentration” that reduces further as the reaction proceeds.

![Figure 9: The concentration of syringic acid and syringaldehyde with time during the electro–oxidation of the lignan at 0.54 V vs Ag/AgCl, pH 12.45.](image)

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† Pre–treatment refers to the dissolution of the oxidizable species in an alkaline solution.
A linear voltammogram of syringic acid showed oxidation peaks at 0.41 V vs Ag/AgCl and 0.60 V vs Ag/AgCl (Figure 10). Since the electro-oxidation of syringaresinol solution occurred at 0.54 V vs Ag/AgCl, any species with an oxidation potential below that value would also be oxidized. Therefore syringic acid underwent some form of oxidation because its first oxidation potential is less than that of 0.54 V vs Ag/AgCl, resulting in a decrease in the concentration of syringic acid (Figure 9).

![Figure 10: Linear voltammogram of syringic acid in a solution of 0.1 M KOH and 0.5 M NaClO₄.H₂O, pH 12.45.](image)

A voltammogram of the organic extract also showed two oxidation potentials. Analysis of the electro-oxidation reaction mixture produced at the lower potential showed that the concentration of vanillin decreased over time whereas the concentration of syringaldehyde increased after approximately four hours (Figure 11).

![Figure 11: The formation of syringaldehyde and vanillin during the electro-oxidation of the organic extract at 0.42 V vs Ag/AgCl, pH 12.45](image)

The percentage concentration of vanillic acid increased very slightly to approximately 0.0078% ($± 0.2048 \times 10^{-2}$%) within the first hour but then decreased about half way through the reaction. The concentration of syringic acid decreased throughout the reaction (Figure 12) because the oxidation potential for this reaction was approximately at the same potential as that of syringic acid (Figure 10).
Figure 12: Formation of syringic acid and vanillic acid during the electro-oxidation of the organic extract at 0.42 V vs Ag/AgCl, pH 12.45

Graphs of the percentage concentration vs time in hours for the above compounds showed that the concentrations of these compounds decreased to values below the starting value during long periods of electro-oxidation. This suggested that the electro-oxidation reactions carried out over long periods of time eventually destroy the compounds that the reaction initially formed. Therefore, a continuous extraction process is required before complete destruction of the products. This was also recommended by Parpot et al. who observed similar results during their studies of vanillin production from the electro-oxidation of lignin obtained via the alkaline Kraft pulping process (Parpot et al., 2000).

Electro-oxidation of the organic extract at the higher oxidation potential produced a maximum concentration of syringaldehyde (0.080% ± 2.606 x 10^{-2}) and vanillin (0.22% ± 8.73 x 10^{-2}) within the first half hour (Figure 13). Thereafter the concentration of both compounds decreased steadily.

The mass of organics present after electro-oxidation was also determined and found to vary (Table 1). A significant reduction occurred particularly during syntheses that were performed at higher oxidation
potentials for the same period of time as syntheses performed at lower potentials suggesting that
electro-oxidation can be used as a means of “cleaning up” the effluent.

Table 1: Masses of organics extracted after each electro–oxidation reaction of the organic extract.

<table>
<thead>
<tr>
<th>Electro–oxidation potentials (E/V vs Ag/AgCl)</th>
<th>Mass of organic extract after electro–oxidation for 1.5 and 5 hours (mg)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1.5 hours</td>
</tr>
<tr>
<td>0.35</td>
<td>39.5</td>
</tr>
<tr>
<td>0.55</td>
<td>47.2</td>
</tr>
</tbody>
</table>

During the electro-oxidation of the concentrated form of the effluent, the percentage concentrations of vanillin and syringaldehyde both increased initially and reached a maximum of 0.0051% (± 0.2581 x 10^-2%) and 0.023% (± 1.757 x 10^-2%) respectively within the first half hour of the reaction (Figure 14). Thereafter, a steady decrease in percentage concentration of syringaldehyde is noted, whereas the percentage concentration of vanillin decreased to a much lesser extent. This again shows that the compounds of interest would need to be extracted from the solution to prevent them from being destroyed.

![Figure 14: Formation of vanillin and syringaldehyde during the electro–oxidation of the concentrated effluent.](image)

The maximum concentration of syringaldehyde (0.011% ± 0.274 x 10^-2%) was reached within the first half hour of the electro-oxidation of lignosulfonates and thereafter decreased. Vanillin also reached a maximum within the first half hour but at a much lower percentage concentration (0.0051% ± 0.4715 x 10^-2%) and thereafter its concentration also decreased (Figure 15).
A literature survey of electro-oxidation reactions carried out on lignin showed the formation of vanillin from the electro-oxidation of alkaline Kraft lignin (Parpot et al., 2000). However, the electro-oxidation reactions had been carried out at 750 A/m² whereas the electro-oxidation reactions in this work were performed at approximately 3 A/m². In order to provide a comparison of the results obtained in this work with that published in literature, the percentage concentration of vanillin obtained was converted to millimolarity (mM) and multiplied 250 times to allow for a theoretical scale up of the current passing through the solution. A result of 0.127 mM was obtained based on the electro-oxidation of a concentrated form of the effluent made up to a solution of 0.15% (m/v) and 0.124 mM of vanillin formed based on the electro-oxidation of 0.15% (m/v) lignosulfonate. These results, using lignin obtained from Sappi's acid bisulfite-based pulping process, compares well with that obtained in literature where a 0.07 mM concentration was obtained based on constant current electrolysis of a 0.5% alkaline Kraft lignin solution (Parpot et al., 2000). This is the first time such investigations of lignin obtained from Sappi's acid bisulfite pulping process have been reported. In addition, this work showed that a maximum amount of vanillin was obtained within half an hour of electro-oxidation compared to 50 minutes as shown in literature for the electro-oxidation of alkaline Kraft lignin. Thus the results obtained for electro-oxidation reactions of lignin from Sappi's effluent based on a theoretical scale up show an increase in the amount of vanillin formed. No previous results on the amounts of syringaldehyde formed from electro-oxidation reactions of acid bisulfite pulping effluent could be found for comparison.

4. Conclusions

The electro-oxidation reactions of the pure compounds syringaldehyde and syringaresinol, show that electro-oxidation reactions of lignin monomers and lignans are possible. Electro-oxidation of syringaldehyde produces its corresponding acid as well as dimers. These dimers, although present in smaller quantities, provide evidence for the polymerization of the lignin monomer which is known to occur in reactions based on radical formation (Pearl, 1952; Pardini et al., 1991; Pearl, 1957; Altwicker, 1967).
Electro–oxidation reactions of the extracted organic component of the effluent, a concentrated form of the effluent and a solution in water of lignosulfonate powder obtained from Lignotech, all showed the formation of syringaldehyde and a smaller amount of vanillin. The results obtained for vanillin compare favorably with that reported in literature; however, the results reported here for syringaldehyde are the first reported on the formation of syringaldehyde obtained from the electro–oxidation of acid bisulfite pulping effluent (Parpot et al., 2000). The compounds of interest must also be removed from the reaction mixture as they form to prevent them being destroyed. Effluent cleaning via electro-oxidation has been shown to be very effective.

The higher concentrations produced at higher potentials were used to calculate the masses of vanillin and syringaldehyde that could possibly be produced after scaling up the reaction 250 times taking into account the flowrates of the effluent to the effluent holding site\(^5\). Table 2 shows large amounts of syringaldehyde and vanillin can be produced by scaling up the reaction concomitant with the large known volumes of 2700 m\(^3\) of effluent produced per day, even though the actual percentage concentrations obtained in this study are low. Higher temperatures should also be considered to further improve the yields of vanillin and syringaldehyde. Further pilot plant studies are needed to determine the industrial feasibility of this study and to optimize the conditions for maximum production of vanillin and syringaldehyde.

**Table 2: Calculated masses of syringaldehyde and vanillin that can be produced from the total effluent produced per day by electrochemical reactions.**

<table>
<thead>
<tr>
<th>Starting Material</th>
<th>Lignan</th>
<th>Organic Extract</th>
<th>Average mass of compounds produced from concentrated effluent and lignosulfonates per day</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Amount produced (kg/day)(^6)</td>
<td>Scaled up (tons/day) (^{**})</td>
<td>Amount produced (kg/day)</td>
</tr>
<tr>
<td>Syringaldehyde</td>
<td>1.12</td>
<td>0.279</td>
<td>6.76</td>
</tr>
<tr>
<td>Vanillin</td>
<td>-</td>
<td>-</td>
<td>19.3</td>
</tr>
</tbody>
</table>

\(^1\) A scale up of 250 times was obtained from a comparison of results for this work with that found in literature\(^5\).

\(^2\) This calculation is based on each batch currently producing 190 m\(^3\) of Ca–spent liquor of which 100 m\(^3\) goes to Lignotech. This leaves 90 m\(^3\) of Ca–spent liquor that goes to the effluent holding site. There are approximately 30 digestions per day resulting in 2700 m\(^3\) of Ca–spent liquor that goes to the effluent holding site per day. These figures were obtained from personnel at Sappi Saiccor.

\(^{**}\) This concentration is based on a typical scale up factor of 250 times obtained from a comparison of the results for this work with that in literature (Parpot et al., 2000)
In addition, the various electro–oxidation reactions carried out on the effluent also show a reduction in the amount of vanillin and syringaldehyde over long periods of electrolysis. Thus electro–oxidation of the effluent can also be used as a means of “cleaning up” the effluent before it is discharged to sea. The preliminary results obtained in this work show a significant reduction in the amounts of the major organics present in the effluent after electro–oxidation of the effluent, and so effective and correct use of this technique could result in a significant reduction of the major organic components present in the total amount of effluent discharged. Studies using electro-oxidation as a means of degrading dyestuffs in water have been performed and have shown significant reductions in color compounds (Mphephu, 1996; Allen et al., 1995). Anodic oxidation reactions are also known to oxidize organic compounds to less harmful products and prolonged electro-oxidation can eventually destroy the compound (Mphephu, 1996; Allen et al., 1995; Simonsson, 1997; El-Ashtoukhy et al., 2009; Patel, 2008; Ugurlu et al., 2008; Ma et al., 2007).

Electro-oxidation has been found to be an environmentally favorable process that does not produce toxic products and is a selective reaction, allowing for easy control in effluent clean up. Overall, the electro–oxidation of Sappi Saiccor’s effluent provides a possible route for the formation of compounds of commercial value without the addition of further chemicals.

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References:


