Improvement of Polypropylene Adhesion by Kraft Lignin Incorporation

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Low surface energy and poor adhesion are well-known characteristics of polypropylene (PP). Surface treatments such as plasma, corona, and laser are usually applied to overcome these limitations. However, current studies highlight the incorporation of hydrophilic or amphiphilic polymers into hydrophobic low-surface-energy polymers as an alternative for increasing surface energy and thus improving adhesion. Lignin could be a promising amphiphilic polymer for use in increasing surface energy. In this work, PP/kraft lignin composites were obtained by incorporating up to 5 wt% of kraft lignin (KL) into a PP matrix. Corona treatment was applied to pristine PP and composites surfaces. Contact angle measurements and peeling tests were carried out to investigate the effects of KL incorporation and corona treatment on the surface energy and the mechanical strength of adhesion. Differential scanning calorimetry (DSC) was used to evaluate the PP’s crystallinity index and recrystallization temperature and to dismiss their effects on the surface energy changes. Scanning electron microscopy (SEM) was applied to investigate the lignin dispersion. The results show that KL incorporation has potential as a method to improve the surface energy of PP, improve its poor adhesion, and enhance the effects of corona treatment.

Keywords: Adhesion, lignin, surface energy, contact angle.

1. Introduction

Polypropylene (PP) can be processed in several ways and is widely used in packaging. However, PP has an apolar and chemically stable surface, which means that it is a hydrophobic material with low surface energy. Consequently, PP is well known for its poor adhesion properties1,2, which can compromise its adhesion to other layers of flexible laminated packaging.

However, several techniques can be employed to improve the adhesion of hydrophobic films, such as Corona3, plasma4, and laser5 treatments. Corona discharge treatment (CDT) has been widely used on PP films, particularly for the possibility of coupling it with extrusion lines and working at room temperature6. CDT consists of applying a non-uniform high-voltage beam from a thin electrode to the surface of the polymeric material, with typical voltages in the range of 10 to 25 kV and frequencies of 10 to 40 kHz7,8. However, for dielectric materials, the technique is mostly limited to films with low thickness since the discharge current decreases drastically as a function of the thickness8. This process oxidizes the polymer surface by adding functional polar groups to the upper layer, including hydroxyl (OH), carbonyl (C=O), and carboxylic (COOH) groups. This mechanism increases the superficial energy of the material, aside from improving the chemical bonding and increasing the roughness, which promote PP adhesion3,8.

Hydrophilization has emerged as an important method to improve the surface energy of hydrophobic polymers. In this process, hydrophilic or amphiphilic materials are blended with hydrophobic materials10. This method of blending different polymers to improve the adhesion of polymer films avoids an additional stages in polymer processing, such as surface treatment, thus reducing the costs of processing and equipment11. In this regard, the lignin appears to be a promising alternative to increase the surface energy of hydrophobic polymers12.

Lignin is a tridimensional amorphous biopolymer that is mainly composed of aromatic moieties that contain different functional groups, such as carboxyl, carbonyl, methoxyl, and hydroxyl13,14. It has been considered as a primary renewable source of aromatics, representing 30% of all non-fossil organic carbon on Earth15,16. Lignin can be obtained from chemical processes such as the kraft process from the pulp and paper industry, where it is treated as waste. Kraft lignin (KL) has a high amount of aromatic hydroxyl groups, which increases its water absorption18. The incorporation of KL into hydrophobic polymers can therefore reduce its hydrophobicity and contribute to the efficiency of methods such as CDT. Furthermore, lignin incorporation into polymeric materials
would be highly beneficial in an economic and environmental sense since it is a readily available, renewable, and non-edible industrial byproduct. Moreover, the application of natural and renewable raw materials on industrial product developments can support the achievement of sustainable societies and contribute to solve problems related to petroleum based products\textsuperscript{19-21}. In addition, composites performance can be improved by proper material selection, which is crucial for achieve sustainable designs and product properties\textsuperscript{22}. In this context, recent methodologies were developed for material selection, which are based on analytical hierarchy process (AHP) and might be applied to solve multi-criteria decision-making (MCDM) problems\textsuperscript{19,20}. These novel methodologies consider evaluation of experts in the field and were successfully applied on natural fiber composites and others\textsuperscript{23-25}.

In this work, up to 5 wt% of KL was incorporated into PP matrix to improve its adhesion properties. Two types of KL were used: acid (AC) and alkaline (ALK) KL, which differ in the amount of hydroxyl level, molar mass, and pH. Typically, AC has a higher hydroxyl level due to purification process applied to reduce the ALK pH. The results indicate that KL has potential as an additive to enhance the adhesion of low-surface-energy polymers and to increase the efficiency of the superficial oxidation of PP by corona treatment.

2. Materials and Methods

PP homopolymer with a density of 0.905 g/cm\textsuperscript{3} and commercially designated as H503HS was supplied by Braskem (Triunfo/Brazil). Technical grade AC and ALK KLS (eucalyptus - hardwood) were kindly supplied by Suzano Papel e Celulose (Suzano/Brazil) and were obtained as byproduct of pulping and paper manufacturing. AC has a pH of 3.5 (2\% ash content), molecular mass of 2.448 g/mol, hydroxyl content of 4.28 mmolOH/g and syringyl (S): guaiacyl (G): p-hydroxyphenyl (H) ratio of 46:50:4. ALK has a pH of 8.2 (2\% ash content), molecular mass of 3.127 g/mol, hydroxyl content of 3.79 mmolOH/g and S:G:H ratio of 47:49:4.

The incorporation of KL into PP was carried out in a melted state using a Brabender Plastograph EC torque rheometer with a W50ET mixer. The processing was done with a cycle time of 8 min at 190 °C, and the lignin was incorporated after 4 minutes (half the time to obtain molten PP). Table 1 shows the sample names and their compositions.

Films with a thickness of 200 µm were obtained in a heated press at 190 °C and pressure of 0.1 MPa for 5 minutes. “CDT” was added to the sample names to identify specimens subjected to corona treatment in addition to the nomenclature presented in Table 1.

2.1 Corona discharge treatment (CDT)

CDT was performed at a fixed velocity of approximately 1.67 m/min with a gap of 3 mm between the film and electrode. The unit energy used in the treatment was 16.8 kJ/m\textsuperscript{2} according to Equation (1):

\[
E = \frac{P}{uL}
\]

Where \(P\) is the discharge power (kJ/s), \(u\) is the linear velocity of the film (m/s) and \(L\) is the length of the discharge electrode (m).

2.2 Contact angle

To evaluate the surface energy of the polymer films, contact angle measurements were performed on the films with AC and ALK addition with and without CDT, and the results were compared to those of neat PP. The measurements were performed by the sessile drop method in a drop shape analyzer DSA100 with two different liquids: distilled water and diiodomethane. The test drop volume was 10 µL, and the measurement was repeated five times for each liquid.

The film surface energy (\(\gamma_{SL}\)) is estimated using the Fowkes model, which was developed from the interfacial free energy model:

\[
\gamma_{SL} = \gamma_s + \gamma_{SL} - 2\sqrt{\gamma_s^d \gamma_{SL}^d} - 2\sqrt{\gamma_s^p \gamma_{SL}^p}
\]

This model combines polar and dispersive interactions between solid and liquid surfaces. \(\gamma_s^d\) and \(\gamma_s^p\) are the polar and dispersive components of surface energy, respectively, while \(\gamma_{SL}^d\) and \(\gamma_{SL}^p\) are the same components for the surface tension of the liquid. Once the liquid surface energy and contact angle are known, the polar and dispersive contribution can be calculated using Equation (3):

\[
\gamma_L (1 + \cos \theta) = 2\sqrt{\gamma_s^d \gamma_L^d} + 2\sqrt{\gamma_s^p \gamma_L^p}
\]

At least two liquids with known surface tension values are required to estimate the polar and dispersive components.
of the total surface energy. Distilled water and diiodomethane were used, and the respective surface tensions for each component are shown in Table 2.

### 2.3 Peel test

Peel tests were performed to evaluate the practical adhesion of neat PP and KL-containing films (with and without CDT) bonded to aluminized bioriented polypropylene (BOPP) layer. The samples were prepared by manual lamination using a polyurethane (PU)-based bicomponent adhesive supplied by Henkel with a 10:8 ratio (stoichiometric condition concerning NCO:OH ratio) of Loctite® Liofol® LA 9526 CP-22 to LA 9729 (polyol and MDI containing prepolymer), which were dissolved in ethyl acetate with a 50% proportion. PP or PP_KL films were bonded to an aluminized surface of BOPP film by PU adhesive with a thickness of 20 µm. The samples were prepared according to the method presented by Ito et. al.

The tests were performed on an EMIC DL-200 peel tester according to ASTM Standard F904. The 90° peel tests were carried out at a speed of 0.280 m/min with a 100-N load cell. The BOPP and PP film edges were secured by clamps in the peel tester, and the tests were performed by pulling a 50-mm length of the PP from the BOPP film at an angle of 90° between the films and the direction of applied force. The test was repeated five times for each condition.

### 2.4 Fourier transform infrared spectroscopy (FTIR)

FTIR analyses were performed to verify the presence of PU adhesive residues in the PP_KL and BOPP films (aluminum surface). The results indicate which type of failure occurred during the adhesive strength test. The analyses were performed using Perkin Elmer Varian 660 equipment in attenuated total reflection (ATR) mode. A total of 16 scans were performed from 4000 to 600 cm⁻¹ with a resolution of 4 cm⁻¹.

### 2.5 Differential scanning calorimetry (DSC)

DSC analyses were performed to evaluate the effect of KL addition on the crystallinity index of PP and the thermal behavior of the blends. Tests were carried out using TA Q200 equipment in accordance with ASTM D3418. An initial heating ramp from 35 to 200 °C was applied to eliminate the material thermal history, followed by a cooling ramp to -50 °C and a third ramp from -50 to 200 °C, where the thermal properties of the samples were evaluated. The analyses were carried out with an approximate mass of 10 mg in an aluminum container. All heating and cooling steps were performed at controlled rates of 10 °C/min and a nitrogen flow of 50 mL/min.

### 3. Results and Discussion

#### 3.1 Free surface energy

Table 3 presents the contact angle measurements. Pure PP presented the highest contact angle measured (i.e., the lowest calculated surface energy; Figure 1). KL incorporation in the PP decreased the contact angles for all tested conditions. The polar contribution to surface energy, which is mainly obtained by the contact angle measured with water, was generally more affected by AC incorporation than ALK addition. Such behavior can be explained by the higher content of polar groups (OH) in the AC than in ALK.

Sample containing 1 wt% AC (without CDT) presented a pronounced increase in surface energy, from 31.93 mN/m to 40.44 mN/m. However, a higher concentration of AC caused a reduction in surface energy, in comparison with PP_AC_1. Incorporation of ALK slightly and stepwise increased the surface energy. ALK (pH 8.2) has a predominant hydrophilic behavior and partial water solubility, which might justify this behavior. Otherwise, despite its higher OH content, AC (pH 3.5) presents amphiphilic behavior and poor water solubility, due to its combination of aromatic and aliphatic bridges and a variety of oxygenated moieties.[17]

CDT-treated samples presented the same trend as untreated specimens, with KL incorporation providing higher surface energy. However, some differences can be pointed out in comparison to non-treated conditions: i) the optimal KL concentration increased to a region close to 2 wt% for both types of KL; ii) ALK generated the highest surface energy value; and iii) the contact angle strongly increased for 5 wt% with both types of KL in comparison to other lignin-containing and corona-treated samples. The results revealed a synergistic effect between KL incorporation and corona treatment, which resulted in higher values of surface energy and could lead to energy savings by using corona treatment due to reductions in time or energy level.

#### 3.2 Adhesive strength

A proper discussion of the peel strength results requires a previous elucidation of the failure type. PU cohesive failure is used to assess the adhesive properties, while adhesive failure is an indirect assessment of adhesive interactions between the substrate and adhesive. Lastly, substrate failure can be used to verify the influence of joint components on the strength.

Figure 2 shows the FTIR spectra from the aluminized surface of BOPP and PP or PP_KL (without CDT treatment) composite surfaces after the peel test, which were previously bonded using PU adhesive. As shown in Figure 2, peaks

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### Table 2. Surface tension components for distilled water and diiodomethane.

<table>
<thead>
<tr>
<th>Surface tension</th>
<th>Water (mN/m)</th>
<th>Diiodomethane (mN/m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total (mN/m)</td>
<td>72.8</td>
<td>50.8</td>
</tr>
<tr>
<td>Polar (mN/m)</td>
<td>51.0</td>
<td>0</td>
</tr>
<tr>
<td>Dispersive (mN/m)</td>
<td>21.8</td>
<td>50.8</td>
</tr>
</tbody>
</table>
assigned to PU groups are present on the aluminized BOPP surface after the peel test. The peaks at 1725 cm$^{-1}$ and 3330 cm$^{-1}$ are related to C=O and NH groups obtained during PU polymerization$^{28}$. The spectra related to PP or PP_KL composite samples do not contain such peaks related to PU groups, which indicates the absence of PU on these surfaces. Thus, the results indicate adhesive failure during the peel test; i.e., PU detached from PP or PP_KL films.

Concerning corona-treated samples, previous work revealed that Al-metalized BOPP layers presented substrate failure in peel tests with loading of 120 N/m or higher$^{26}$. Table 4 presents the average peel strength values. Corona-treated samples provided average peel strength higher than 126.12 N/m, which suggest that Al substrate failure of these peel specimens as well according to Ito et. al$^{26}$. KL-containing samples (without CDT) developed higher peel strength, in comparison with neat PP. No significant difference was observed between both types of KL regarding peel strength when taking the error involved in the experiments into consideration. KL addition increased the peeling strength, which is in agreement with contact angle measurements since KL incorporation also increased the surface energy, and adhesive failure was verified. There was some divergence between the surface energy and peel strength evolution as a function of KL content. This can be explained by the limitations of the contact angle measurement and peel test, which can only assess the adhesion strength indirectly.

Since Al cohesive failure was assumed, the peel results for CDT-treated samples are related to the KL incorporation effect on the Al layer strength. KL addition also increased the peel strength of CDT-treated adhesively bonded joints. A concentration of 1 w% KL generated the highest peel strength values. KL incorporation could thus be a tool for improving CDT efficiency when applied to PP surfaces, which would reduce the required corona energy level or obtain higher surface energy for the same discharge level.

### 3.3 Differential scanning calorimetry (DSC)

The polymer crystallization index or degree strongly influences thermoplastic adhesion, in which adhesive interactions are usually developed by the amorphous phase of such polymers. In this context, a decrease in the degree of crystallization by KL incorporation could contribute to the increased peel strength. DSC was used to investigate the effect of lignin incorporation on the melting temperature ($T_m$), recrystallization temperature ($T_c$), glass transition temperature ($T_g$), and degree of crystallization ($\chi_c$%). The results are shown in Table 5. The degree of crystallinity of PP was calculated according to Equation (4):

$$X_c(\%) = \frac{\Delta H_m}{\Delta H^m_{pp}} \times \frac{1}{f} \times 100$$  \hspace{1cm} (4)

Where $\chi_c$ is the degree of crystallinity (%), $\Delta H_m$ is the enthalpy of fusion of different blends, $\Delta H^m_{pp}$ is the enthalpy of fusion of 100% crystalline PP (207 J/g)$^{29}$, and $f$ is the PP fraction in blends composition.

Figure 3 shows the thermograms of the second heating and cooling cycles for PP and PP_KL. The addition of lignin shifts and slightly broadens both the melting and crystallization peaks. Only one glass transition temperature was found for PP_KL. It is well established that the presence of only one $T_g$ is considered to indicate homogeneity at the segmental level and therefore blend miscibility$^{30}$. However, this general rule cannot be correctly applied to this system. $T_g$ of lignin is usually within the range of 110-160 ºC$^{31}$.

### Table 3. Contact angle values using water and diiodomethane.

<table>
<thead>
<tr>
<th>Material</th>
<th>Contact angle (º)</th>
<th>Material</th>
<th>Contact angle (º)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Water</td>
<td>Diiodomethane</td>
<td>Water</td>
</tr>
<tr>
<td>PP</td>
<td>107.22±2.70</td>
<td>56.70±0.96</td>
<td>PP_CDT</td>
</tr>
<tr>
<td>PP_AC_1</td>
<td>91.42±1.34</td>
<td>48.88±1.74</td>
<td>PP_AC_1_CDT</td>
</tr>
<tr>
<td>PP_AC_2</td>
<td>92.42±0.96</td>
<td>51.80±1.25</td>
<td>PP_AC_2_CDT</td>
</tr>
<tr>
<td>PP_AC_5</td>
<td>92.46±4.47</td>
<td>51.84±1.83</td>
<td>PP_AC_5_CDT</td>
</tr>
<tr>
<td>PP_ALK_1</td>
<td>97.86±1.26</td>
<td>56.80±0.70</td>
<td>PP_ALK_1_CDT</td>
</tr>
<tr>
<td>PP_ALK_2</td>
<td>96.42±0.86</td>
<td>56.00±0.96</td>
<td>PP_ALK_2_CDT</td>
</tr>
<tr>
<td>PP_ALK_5</td>
<td>95.58±1.47</td>
<td>54.34±0.75</td>
<td>PP_ALK_5_CDT</td>
</tr>
</tbody>
</table>

Figure 1. Surface free energy as a function of lignin concentration (AC and ALK) with and without surface treatment.
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Figure 2. FTIR spectra for PP films with (a) AC and (b) ALK (without CDT) compared to BOPP after peel strength. Amplifications of the bands of 1725 cm⁻¹ (c and d) and 3330 cm⁻¹ (e and f) for AC and ALK, respectively.

Table 4. Peel strength values for PP and PP KL films.

<table>
<thead>
<tr>
<th></th>
<th>Average Strength (N/m)</th>
<th>Average Strength (N/m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PP</td>
<td>8.73 ±0.84</td>
<td>PP_CDT</td>
</tr>
<tr>
<td>PP_AC_1</td>
<td>10.61 ±0.57</td>
<td>PP_AC_1_CDT</td>
</tr>
<tr>
<td>PP_AC_2</td>
<td>8.91 ±0.91</td>
<td>PP_AC_2_CDT</td>
</tr>
<tr>
<td>PP_AC_5</td>
<td>12.67 ±1.95</td>
<td>PP_AC_5_CDT</td>
</tr>
<tr>
<td>PP_ALK_1</td>
<td>10.50 ±1.28</td>
<td>PP_ALK_1_CDT</td>
</tr>
<tr>
<td>PP_ALK_2</td>
<td>9.38 ±1.41</td>
<td>PP_ALK_2_CDT</td>
</tr>
<tr>
<td>PP_ALK_5</td>
<td>9.08 ±0.99</td>
<td>PP_ALK_5_CDT</td>
</tr>
</tbody>
</table>
Moreover, lignin stiff molecule hinders $T_g$ measurements when blended or diluted in other polymers. Therefore, the $T_g$ associated with the lignin phase of PP_KL could not be determined, which was probably due to the overlap of $T_g$ of KL with the large PP melt endotherm.

The degree of crystallization was increased by the incorporation of both types of KL at concentrations of up to 2 wt%. However, the highest KL content presented a slight decrease in the degree of crystallization. Lignin might act as heterogeneous nucleating agent in thermoplastic polymers. The nucleating agent effect of 1% lignin incorporation has been demonstrated for poly(3-hydroxybutyrate) (PHB). Similar to this study, blends of PP and lignin have previously exhibited an increase in crystallinity upon lignin incorporation. These findings suggest that lignin favors crystallization at small amounts, but at higher content, its amorphous structure overcomes the heterogeneous nucleation effect, and a lower degree of crystallinity is achieved. Nevertheless, no meaningful changes on crystallinity degree were found for all lignin contents, as listed in Table 5. In addition, higher degrees of crystallization should reduce the peel strength due to the decrease in the amorphous phase content, and this correlation is not observed when comparing DSC and peel strength results (Table 4 and Table 5). Therefore, the crystallinity degree cannot be considered to influence the peel strength results, which strengths our arguments that there is an increase in surface energy due to lignin addition. Moreover, the investigation of the thermal behavior of PP/lignin blends are of uttermost importance for further technological applications.

### 3.4 Scanning electron microscopy (SEM)

Scanning electron micrographs were obtained to determine whether phase separation occurred between KL and PP phases for the KL contents tested. A lignin rich phase, as consequence of KL aggregation or phase separation, might act as stress concentration factor and low cohesive strength regions, which could result into early failure of flexible laminate. Figure 4 shows the micrographs of AC_KL and a cut surface of PP_AC. The micrographs revealed an absence of patterns related to phase separation. As shown in Figure 4a, AC_KL presented an irregular colloidal shape, which cannot be identified in AC-containing PP. Besides the well-known immiscibility between PP and KL, the results pointed out an absence of KL phase, which demonstrates good dispersion between the phases. The same behavior was verified for ALK-containing samples. Results are in agreements with previous works, in which lignin aggregation were verified for concentration of 20 wt% into PP, or 10% into PBAT.

<table>
<thead>
<tr>
<th></th>
<th>Tm (°C)</th>
<th>Tg (°C)</th>
<th>Tc (°C)</th>
<th>Crystallinity Degree (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PP</td>
<td>164.58</td>
<td>-5.69</td>
<td>115.30</td>
<td>50.77</td>
</tr>
<tr>
<td>PP_AC_1</td>
<td>166.00</td>
<td>-3.16</td>
<td>115.26</td>
<td>51.92</td>
</tr>
<tr>
<td>PP_AC_2</td>
<td>165.44</td>
<td>-3.96</td>
<td>114.32</td>
<td>51.37</td>
</tr>
<tr>
<td>PP_AC_5</td>
<td>166.32</td>
<td>-3.70</td>
<td>114.18</td>
<td>49.02</td>
</tr>
<tr>
<td>PP_ALK_1</td>
<td>166.71</td>
<td>-5.04</td>
<td>115.24</td>
<td>51.53</td>
</tr>
<tr>
<td>PP_ALK_2</td>
<td>166.53</td>
<td>-5.65</td>
<td>114.78</td>
<td>51.71</td>
</tr>
<tr>
<td>PP_ALK_5</td>
<td>166.88</td>
<td>-3.61</td>
<td>114.14</td>
<td>48.31</td>
</tr>
</tbody>
</table>
4. Conclusions

PP_KL composites were obtained, and the effects of KL incorporation on practical adhesion were investigated. KL incorporation increased the surface energy of pristine and CDT-treated samples. CDT-treated samples showed the highest surface energy values with 2 wt% of KL, while the highest peeling strength was obtained for specimens with 1 wt% of KL. A synergistic effect of KL and CDT was observed. The surface energy changes are not related to changes in the PP crystallinity index. PP and KL phase separation was not observed in SEM micrographs. KL addition improved the practical adhesion of PP films and increased the efficiency of CDT.

5. Acknowledgements

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6. References


