Comparison of thermal behavior of natural and hot-washed sisal fibers based on their main components: cellulose, xylan and lignin. TG-FTIR analysis of volatile products

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Abstract:

This paper presents in a comprehensive way the thermal behavior of natural and hot-washed sisal fibers, based on the fundamental components of lignocellulosic materials: cellulose, xylan and lignin. The research highlights the influence exerted on the thermal stability of sisal fibers by other constituents such as non-cellulosic polysaccharides (NCP) and mineral matter.

Thermal changes were investigated by thermal X-ray diffraction (TXRD), analyzing the crystallinity index (%Ic) of cellulosic samples, and by simultaneous thermogravimetric and differential thermal analysis coupled with Fourier-transformed infrared spectrometry (TG/DTA-FTIR), which allowed to examine the evolution of the main volatile compounds evolved during the degradation under inert and oxidizing atmospheres. The work demonstrates the potential of this technique to elucidate different steps during the thermal decomposition of sisal, providing extensible results to other lignocellulosic fibers, through the analysis of the evolution of CO₂, CO, H₂O, CH₄, acetic acid, formic acid, methanol, formaldehyde and 2-butanone, and comparing it with the volatile products from pyrolysis of the biomass components. The hydroxyacetaldehyde detected during pyrolysis of sisal is indicative of an alternative route to that of levoglucosan, generated during cellulose pyrolysis.

Hot-washing at 75 °C mostly extracts non-cellulosic components of low decomposition temperature, and reduces the range of temperature in which sisal decomposition occurs, causing a retard in the pyrolysis stage and increasing Tₘₐₕₐ₉ₙ and Tₜₖₖₖₕₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖₖ₆

Keywords: Sisal fiber; Hot-water treatment; Biomass components; TGA-FTIR; Gas evolution; Pyrolysis; Combustion.
1. Introduction

The large number and diversity of lignocellulosic products as well as plurality of their applications in many industrial sectors, implies the need to comprehensively know their thermal behavior, stability and degradation processes at elevated temperatures.

Pyrolysis studies of synthetic biomass, obtained by mixing its fundamental constituents, revealed that, in general, the overall degradation of biomass follows the principle of additivity [1,2]. However, different authors [3,4] show that interaction among the constituents exits, whereby the proportion of the pyrolysis products differs from that of natural biomass.

It has been demonstrated that the addition of extractives and mineral compounds to the fundamental biomass components [5-7], and the degree of crystallinity of cellulose [8], affect the pyrolysis process. The influence of extractives and mineral matter on thermal decomposition of different lignocellulosic materials has been studied extensively [9-11]. Water washing is a simple and economical pretreatment, which has been used to remove alkali metals from biomass, to avoid the toxicity of mists generated by the combustion of biomass, so preventing accumulation problems (deposition, corrosion, erosion, etc.) in combustion and gasification plants, and also to improve process yields [12-14]. Hot washing process reduces the proportion of substances adhered to the cell walls, removing organic compounds, as a hemicellulose fraction and other low molecular weight carbohydrates, and minerals such as salts of alkali, alkaline earth and transition metal cations.

In addition to the traditional purposes, the use of Agave species fibers has spread to a variety of processes of interest in many industrial sectors, due to their ease of cultivation, with high productivity and low cost associated, as well as their renewable and biodegradable character. Furthermore, their biomass component proportion makes them suitable for processability and biofuel production [15,16]. In particular, Agave sisalana fiber or sisal fiber is one of the most widely used due to its good mechanical properties, used in a great variety of traditional applications due to its hardness, coarseness and resistance to wear, which has gained great interest during the last decades in the manufacturing of composite materials, particularly as reinforcement of mortars, concrete and polymeric matrices [17-19], as polypropylene [20-22].

Sisal fibers are mainly composed of cellulose and non-cellulosic polysaccharides (NCP): hemicellulose O-acetyl-(4-O-methylglucurono)xylan [23] and pectins composed of rhamnose, arabinose and uronic acid at a 2.3:1:3.3 ratio [24], along with a smaller amount of lignin, whose molar ratio of constitutional units, p-hydroxyl phenol (H), guaiacyl (G) and syringyl (S), has been established in 2% H, 22% G and 76% S [25]. Furthermore, the fiber is constituted by a lower proportion of extractives, such as lipophilic components [26], simple phenolic derivatives and mineral matter, mainly represented by calcium oxalate deposits located between the elementary fibers comprising the fiber bundle [27]. The composition of sisal, as any natural fiber, varies depending on the origin, age and extraction process of the components, among other factors. The compositional variability in terms of constituents is very large as can be deduced from different reviews [15,28,29].

There is a lot of information scattered in the literature about the thermal behavior of sisal fibers, but few studies have analyzed it in depth [30-32]. The extensive work recently published by Martin et al.
[33] focuses on the thermal decomposition of sisal and the main biocomponents extracted from the fiber itself. However, given the complexity of thermal processes occurring in lignocellulosic materials, the comparison of thermogravimetric stages with their associated energy effects requires a detailed analysis, interpreted through the analysis of the decomposition of pure biomass components, adding a new perspective that would allow for a better definition of the phenomena occurring. Furthermore, the evolution of the thermal degradation products of sisal fibers is poorly described. Only the liquid fraction released during fast pyrolysis has been characterized [34], there being a lack of information about the gaseous products emitted.

The thermogravimetry coupled with Fourier-transformed infrared spectroscopy (TG-FTIR) technique has been proved to be a valuable and simple tool, not only to analyze the gas phase during the thermolysis of lignocellulosic compounds [35-37], but also to clarify different steps during fiber decomposition.

This paper aims to analyze and present in a comprehensive way the thermal behavior of natural and hot washed sisal fibers, discussing their thermogravimetric and differential thermal analysis (TG-DTA) on the basis of the degradation of fundamental components, cellulose, lignin and xylan, and comparing the evolution of the gas phase decomposition products by TG-FTIR analysis.

2. Materials and Methods

The sisal fibers used in this work were provided by Cayetano García del Moral S. L. (Cabra de Santo Cristo, Jaén, Spain). They were cut to 2 - 4 mm length and then subjected to a washing process with stirring in hot water at 75 °C for 2 hours. Hereinafter the samples will be referred to as Natural Sisal and Washed Sisal. Microcrystalline cellulose powder (Aldrich), alkali low sulfonate content lignin (Sigma Aldrich) and birch xylan (xylose ≥ 90%, Aldrich) were used as representatives of the main biomass components.

X-ray diffraction (XRD) measurements were performed in a Philips X’Pert PRO MPD, with a X’Celerator real time multiple strip RTMS system. Thermodiffraction was performed in Helium atmosphere at temperatures up to 400 °C, using a TTK450N chamber for Aldrich cellulose and a one HTK1200N for Natural Sisal. Spectra were acquired every 20 °C, each one was recorded for 50 minutes in the 2θ range from 5 to 45°. Temperature was linearly changed between isotherms at 5 ºC/min, and the stabilization time at each isotherm was 5 minute. Data were analyzed employing X’Pert HighScore software. The crystallinity index, %Ic(XRD), was calculated at several temperatures using the equation proposed by Segal [38]:

\[
\%I_c(XRD) = 1 - \frac{I_{am}}{I_{max}} \times 100
\]

where \(I_{max}\) is the maximum intensity of the (002) lattice reflection of cellulose Ib type, located at 2θ between 22 and 23°, and \(I_{am}\) is the intensity attributed to amorphous part, given at 2θ between 18° and 19° for Ib cellulose.
Thermogravimetric study was performed in a TA Instruments STD 2960 simultaneous thermal analyzer. The experimental setup consisted of a 10 °C/min linear heating program from 20 to 750 °C, with flow rates of 50 and 120 mL/min, both in air and in nitrogen. The gases released were immediately analyzed in a Bruker VECTOR 22 infrared spectrometer, coupled to the outlet port of the SDT. The transfer line and the cell were kept at a temperature of 200 °C. The spectrum of the evolved gas was recorded every 61 seconds, and consists of an average of 64 scans with a resolution of 4 cm⁻¹ in the range from 4000 to 500 cm⁻¹. Spectra were analyzed using Bruker OPUS 5.5 software, being baseline-corrected by a concave rubber band algorithm computed with 10 iterations on 64 points.

Ashes from the combustion of natural and washed sisal fibers were semi quantitatively analyzed by X-ray fluorescence, using a PANalytical AXIOS spectrometer.

3. Results and Discussion

3.1. Evolution of the crystallinity index of cellulose and sisal

The diffraction patterns at different temperatures of microcrystalline cellulose and Natural Sisal are shown in Fig. 1. At room temperature, cellulose clearly presents reflections around 2θ = 15, 16.5, 20.5, 22.5 and 34.5°, corresponding respectively to the planes (101), (10\overline{1}), (021), (002) and (040) of I₉ type allomorph, while sisal fiber only exhibits the main (002) peak, and a convolution of (101)-(10\overline{1}) reflections, which are superimposed to calcium oxalate hydrate Ca(COO)₂·H₂O ones (pattern 20-0231 of the Joint Committee on Powder Diffraction Standards - International Center for Diffraction Data, JCPDS – ICDD). For both, it is observed that the intensity of reflections decreases with increasing temperature, shifting the diffraction peaks towards a greater spacing. It can be explained due to the progressive deterioration of cellulose crystal lattice, until the structure becomes completely amorphous, observed before 310 °C. Furthermore, the constitutive minerals of sisal fiber experiment transformations with increasing temperature, generating stable crystalline compounds under inert atmosphere.

![Figure 1. X-ray diffractograms at different temperatures, a of microcrystalline cellulose and b of Natural sisal, where can be observed reflexions from Ca(COO)₂·H₂O (■).](image-url)
The evolution of %\textit{I}\textsubscript{c}(XRD) with temperature for both cellulose and sisal is displayed in \textbf{Fig. 2}, being observed a much more pronounced variation for the natural fiber. A rapid decrease occurs for sisal index at temperatures below than 100 °C, remaining stable at around 42% until 210 °C. In contrast, it diminishes slowly to 76% for cellulose. Near 230 °C, the crystallinity index of both sisal and cellulose falls sharply, presenting almost the same slope in both cases. At 270 °C it reaches the smallest value of 17% for sisal, and 65% for the studied cellulose, which is totally amorphized above 310 °C. This behavior for sisal fiber is close to that reported by Saikia [32], which describes a reduction of the degree of crystallinity from 51 to 41% when sisal is heated in air at 177 °C, becoming completely amorphous at around 260 °C.

\textbf{Figure 2.} Variation of the crystallinity index, %\textit{I}\textsubscript{c}(XRD), with temperature for microcrystalline cellulose (▲) and Natural Sisal (●).

3.2. TG-DTG/DTA analysis of the biomass components

Previously to the analysis of the thermal decomposition of sisal fibers, it is important to understand the behavior of the main lignocellulosic components. \textbf{Figure 3} exhibits TG, DTG and DTA plots for the studied cellulose, xylan, and lignin, under both air and nitrogen atmospheres. Temperature ranges, temperature of maximum mass loss rate (Tb) and the mass loss on a dry basis are summarized in \textbf{Table 1}. The weight on dry basis is calculated from each thermogram excluding the mass loss below 180°C.
Figure 3. TG/DTG-DTA thermograms of the main biomass components, under nitrogen (a-b) and air purges (c-d) with flow rates of 120 mL/min. The inner inset in Fig. 3b notes three superimposed stages on DTG curve of lignin.

Table 1: Thermoanalytical data TG/DTG-DTA of biomass components, under air and nitrogen atmospheres (120 mL/min flow rate).

<table>
<thead>
<tr>
<th></th>
<th>Air</th>
<th>Nitrogen</th>
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<tbody>
<tr>
<td></td>
<td>$T_b$ (°C)</td>
<td>% wt $^b$</td>
</tr>
<tr>
<td><strong>Cellulose</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>21 – 185</td>
<td>65</td>
<td>52 endo</td>
</tr>
<tr>
<td>185 – 329</td>
<td>325</td>
<td>325 endo</td>
</tr>
<tr>
<td>329 – 354</td>
<td>333</td>
<td>17.9</td>
</tr>
<tr>
<td>354 – 737</td>
<td>473</td>
<td>19.9</td>
</tr>
<tr>
<td>Ash (737 ºC)</td>
<td>0.3</td>
<td></td>
</tr>
<tr>
<td><strong>Xylan</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>20 – 165</td>
<td>88</td>
<td>90 endo</td>
</tr>
<tr>
<td>165 – 298</td>
<td>247</td>
<td>58.9</td>
</tr>
<tr>
<td>298 – 423</td>
<td>344</td>
<td>31.2</td>
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<tr>
<td>423 – 502</td>
<td>435</td>
<td>1.3</td>
</tr>
<tr>
<td>502 – 653</td>
<td>562</td>
<td>0.7</td>
</tr>
<tr>
<td>Ash (737 °C)</td>
<td>7.3</td>
<td></td>
</tr>
<tr>
<td><strong>Lignin</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>26 – 180</td>
<td>86</td>
<td>98 endo</td>
</tr>
<tr>
<td>180 – 344</td>
<td>305</td>
<td>22.8</td>
</tr>
<tr>
<td>344 – 426</td>
<td>380</td>
<td>6.6</td>
</tr>
<tr>
<td>426 – 737</td>
<td>456</td>
<td>8.1</td>
</tr>
<tr>
<td>Ash (737 °C)</td>
<td>62.5</td>
<td></td>
</tr>
</tbody>
</table>

$^a$ DTG peak temperature

$^b$ mass loss percentage and ash/residue percentage in dry base
It can be distinguished fundamentally three stages, described underneath.

**i)** In the low temperature range, below 180 °C, fundamentally physically adsorbed water evaporates. The mass losses account for about 4% for cellulose, 7% for xylan and 9% for lignin. The process implies a large endothermic effect with DTA peaks under nitrogen atmosphere at 56, 80 and 90 °C, respectively.

**ii)** Between 200 and 400 °C processes of depolymerization, decomposition and rearrangement occur, involving a considerable mass loss. Chemical bonds of the starting biomass break (primary reaction), generating a reactive residue (reactive char) plus liquid (tar) and gaseous compounds, taking place under inert and air atmosphere by different mechanisms, pyrolysis and oxidative pyrolysis respectively. The last one explains an accelerated decomposition in the presence of oxygen, related to oxidative type reactions described for cellulose [39,40] and for lignin [41], which in turn results in the advancement of the maximum rate of decomposition. At this stage, endothermal volatilization and exothermal charring processes are responsible of the resulting energy balance [42,43].

Comparing biomass components, xylan decomposes at lower temperatures, followed by cellulose, while lignin does it over a wider range of temperatures, due to its heterogeneous structure, comprising aromatic rings with multiple branches and functional groups, whose bonds cleave along a wide range of temperature.

Two sub-stages of exothermic nature can be observed in hemicellulose decomposition. The first one, very slight with Tb at around 228 ºC, can be associated to the decomposition of residual side chains, still present in the commercial xylan, assigned to the decomposition of 4-O-methyl glucuronic acid and acetyl groups of xylan by Shen et al. [44], while the second sub-stage, located at around 286 ºC, is due to rupture of the xylan main chain. These sub-stages are hardly distinguishable in air, where the decomposition process occurs at much higher speed than in inert atmosphere.

Cellulose is the biomass component which presents the maximum mass loss and maximum mass loss rate. Pyrolysis of cellulose can be described in a simplified form as a mechanism of chain depolymerization via transglycosylation reactions [45]. One important endothermic process is observed in nitrogen atmosphere at Tb = 338 °C, associated to the heat of evaporation of volatile products generated by the pyrolysis of the glycosyl units [40], and it matches with the collapse of the structure and the complete loss of the crystalline structure, as shown in **Fig. 1A**. In contrast, a sharper endotherm at Tb = 325 °C is observed in air. This shifting to lower temperature is due to the oxidative degradation of cellulose which starts at lower temperatures as a result of radical autoxidation mechanism that occurs in the presence of oxygen [39]. In air, immediately after the endothermic process, it can be distinguished a large exothermic one with Tb at 333 ºC and maximum DTA peak at 350 ºC, associated to the oxidation of volatiles generated in the preceding endothermic process.

Lignin is the one of the biomass components which experiences the lower mass loss and at the lowest rate. Its thermal decomposition, in which at least three superimposed processes can be distinguished at around 278, 315 and 358 ºC in nitrogen atmosphere (as shown the inset of the **Fig. 3B**), present an overall exothermic process of greater intensity in air than in nitrogen. Furthermore, after pyrolysis stage, lignin is the component which presents the highest char content, about a 60%, generated
by condensation of the phenolic groups during its thermal degradation under inert atmosphere. The G-type lignin used in this work probably contributes to a higher char generation due to the own nature of the biomass polymer that, in one hand, allows a high degree of crosslinking facilitated by a reduced content in methoxyl groups, which in turn increases its thermal stability, and that, on the other hand, due to the own constituent guaiacyl units, is prone to experience thermal condensation reactions [46].

iii) At temperatures higher than 350-400 °C, the thermal behavior depends on the atmosphere. Under inert gas, the structure of the products generated in the previous stage starts to change through cleavage of bonds and volatilization of heteroatoms, involved in the condensation and reorganization processes leading to the formation of products with high carbon content. This carbonization stage is accompanied by a small and gradual mass loss and by slight energy changes, as it can be respectively observed on the TG and DTA plots. In addition, the volatilized compounds undergo secondary thermolytic reactions in gas phase, which do not affect the mass of the remaining sample. The effect on the DTA signal is weak since the thermocouples of the instrument are located at the bottom of the sample and reference crucibles, relatively distant from where the evolved gases may react. Fisher et al. [47] maintain that secondary reactions can be thermally neutral, depending on the conditions under which the char is generated. In the case of cellulose, a small exothermic effect is observed next to the 338°C endotherm, at around 400 °C, which is associated to the charring process. Milosavljevic et al. [48] claim that the charring process of the cellulose can be exothermic (-2.6 kJ/g referred to residual carbon), endothermic (0.86 kJ/g) or combination of both, depending on the products generated. Similarly, exothermic effects at around 460 °C in the case of xylan, and at 480 and 650 °C in the case of lignin, are associated to carbonizing processes.

In the presence of oxygen, the oxidation of volatile products (homogeneous reaction) and combustion of the char (heterogeneous reaction) take place in the range between 350 and 500 °C, depending on the nature of the lignocellulosic material. These exothermic phenomena appear at lower temperature and with higher intensity in xylan (390 ºC) than in cellulose (475 ºC). In addition, small exothermic effects are observed at higher temperatures, which can be associated to residual combustion processes, accompanied by a very small mass loss, which will be discussed afterwards. However, in the case of lignin, the combustion stage does not constitute a noticeable phenomenon. It appears to pass through several consecutive mild stages, with small mass loss at Tb = 380 and 456 °C. Moreover, unlike what may be expected, the percentage of the final residue of lignin is slightly greater in air than in nitrogen (63% > 58%). This can also be explained by several reasons: a) that a partial oxidation has occurred, generating compounds of higher molecular weight; b) that a stable char has been formed due to the elevated content in aromatics [49]; c) that the metals present in lignin oxidize, thus increasing their mass. Di Blasi [50] describes many factors that affect the reactivity of the char which could explain the elevated percentage that remain after thermal decomposition of lignin under air.

The data presented here confirm that the thermal stability of the biomass components decreases in the following order: lignin, cellulose and hemicellulose. These different behaviors are due to their different chemical composition and structure [51]. Hemicellulose has an amorphous structure, formed by
various polysaccharides with branches of different compositions, while cellulose is a polymer composed of linear chains of glucose, with a longer range order and a degree of polymerization of ten to one hundred times higher than that of hemicellulose. However, lignin is a highly cross-linked polymer consisting of highly branched units of phenylpropane, which confers a great thermal stability.

3.3. TG-DTG/DTA analysis of natural and hot-washed sisal fibers

Thermal degradation of sisal and washed sisal has been studied by means of simultaneous TG-DTA using purges of air and nitrogen, both at 50 and 120 mL/min. Figure 4 shows a comparison of their thermogravimetric curves in air and nitrogen. Table 2 shows the mass loss on a dry basis, ranges of temperature and maximum temperature of the mass loss rate (Tb) of each stage.

Figure 4. Comparison of TG/DTG-DTA thermograms of Natural and Washed Sisal, under 50 mL/min flow rate.
Table 2 Thermoanalytical data TG/DTG-DTA of Natural and Washed Sisal, under air and nitrogen.

<table>
<thead>
<tr>
<th></th>
<th>Air</th>
<th>Nitrogen</th>
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<tbody>
<tr>
<td></td>
<td>$T_{b^a}(^\circ C)$</td>
<td>% wt $^b$</td>
</tr>
<tr>
<td>Raw sisal</td>
<td>20 – 186</td>
<td>48</td>
</tr>
<tr>
<td>50 mL/min</td>
<td>186 – 294 (203,281)</td>
<td>21.1</td>
</tr>
<tr>
<td></td>
<td>294 – 380</td>
<td>336</td>
</tr>
<tr>
<td></td>
<td>380 – 487</td>
<td>435 (47)</td>
</tr>
<tr>
<td>Ash ($737^\circ C$)</td>
<td>1.2</td>
<td></td>
</tr>
</tbody>
</table>

|                    | $T_{b^a}(^\circ C)$ | % wt $^b$ | $T_{p_{\text{peak}}}(^\circ C)$ $^c_{\text{DSC}}$ | $T_{b^a}(^\circ C)$ | % wt $^b$ | $T_{p_{\text{peak}}}(^\circ C)$ $^c_{\text{DSC}}$ |
|--------------------| 24 – 187             | 62     | 53 endo                                      | 20 – 217              | 62      | 61 endo                                      |
| Washed sisal       | 187 – 301             | 294     | 17.0 352 exo                                 | 171 – 314             | 293     | 18.5 293 exo                                 |
| 50 mL/min          | 301 – 391             | 329     | 60.0 457 exo                                 | 314 – 474             | 352     | 63.6 355 exo                                 |
|                    | 391 – 458             | 442     | 21.1 480 exo                                 | 474 – 737             | 651     | 651 exo                                      |
| Ash ($737^\circ C$)|                            0.3   |                          | Residue ($737^\circ C$)   | 11.2                                      |
| Washed sisal       | 26 – 173              | 66     | 58 endo                                      | 23 – 170              | 70      | 55 endo                                      |
| 120 mL/min         | 173 – 301             | 295     | 17.6 351 exo                                 | 170 – 311             | 294     | 17.9 294 exo                                 |
|                    | 301 – 381             | 328     | 58.5 448 exo                                 | 311 – 379             | 350     | 59.4 355 exo                                 |
|                    | 381 – 451             | 432     | 20.9 475 exo                                 | 379 – 737             | 637     | 637 exo                                      |
| Ash ($737^\circ C$)|                            0.8   |                          | Residue ($737^\circ C$)   | 11.4                                      |

$^a$ DTG peak temperature  
$^b$ mass loss percentage and ash/residue percentage in dry base

Three main stages can be distinguished during thermal decomposition of sisal fiber:

**i)** Evaporation of physically adsorbed water mainly occurs at temperatures below 100 °C. The process represents a mass loss of 4 - 9% and is accompanied by an endothermic effect. In this range of temperature, an important decrease of crystallinity is observed in sisal (Fig. 2).

**ii)** Both with air and with nitrogen, two sub-stages are observed in the 200-400 °C range of thermal decomposition of sisal fiber.

**ii.1)** The first sub-stage, below 300 °C, is attributed mainly to the degradation of non-cellulosic polysaccharides (NCP) of sisal, both of hemicellulose and pectins. Decomposition of sisal lignin also takes place, but in a lesser extent because of its low proportion. A mass loss of 17 - 21% is observed in this sub-stage for natural sisal, being slightly lower for washed sisal. The temperature at maximum mass loss rate, $T_{b_{\text{NCP}}}$, is around 280 °C for natural sisal and 294 °C for washed sisal, both in air and inert atmosphere, which are close to that of xylan decomposition, discussed in the previous section. The DTA plot obtained from washed sisal in nitrogen shows an exotherm with peak at 294 °C, also observed in air at 293 °C. This sub-stage coincides with the complete amorphization of the sisal cellulose, observed in Fig. 1B at temperatures above 290 °C. The accomplished analysis presents a clear correspondence
between DTA and thermogravimetric changes. The observed values are slightly different from those reported by Martin et al. [33], who described the maximum mass loss rate temperature at about 20 °C above the values found in this work.

ii.2) The second sub-stage, between 300 and 400 °C under nitrogen, corresponds almost exclusively to the degradation of cellulose by breaking of glycosidic linkages of the glucose chain. Pyrolysis of lignin also continues, although to a much lower extent. In air atmosphere, the combustion of NCP superimpose to the decomposition of sisal cellulose. This sub-stage shows a mass loss of almost 50% in the case of natural sisal, and about 60% for washed sisal. In air, $T_{b,CEL}$ is in the range from 336 to 339 °C for natural sisal and between 328 and 332 °C for washed sisal, while in nitrogen the differences shorten, falling in the range from 348 to 353 °C. Martin et al. [33] reported the same trend for the raw and defatted sisal, but at temperatures 20 to 30 ºC higher than those found in this study. The degradation temperature of cellulose in sisal is slightly higher than that of pure cellulose, discussed in the previous section. It can be attributed to the protective effect of the rest of components present in sisal. The analysis in an inert atmosphere reveals that this step is accompanied by an endothermic effect, at about 355 °C, due to the breaking of bonds, as it happens in the pure cellulose. However, in air an intense exotherm is observed at around 350 °C, associated to the oxidation of the volatiles generated. But differently than cellulose, the endotherm preceding the exotherm is not observed. It is probably hidden by the exotherm resulting from combustion of sisal hemicellulose and other NCP.

It must be noted that the mass loss of the second sub-step is slightly higher in nitrogen atmosphere than in air (Table 2). The oxygen gain that saccharides undergo under oxidizing atmosphere can explain this difference, as Saikia concluded [32]. This phenomenon is also observed, even more pronounced, in biomass pure compounds.

iii) The third stage starts at 400 °C. A small and sustained mass loss is observed in inert atmosphere, which corresponds to the process of carbonization. Very mild exothermic processes are observed in washed sisal, at around 650 and 635 °C, with gas flows of 50 and 120 mL/min, respectively, but no substantial change in mass. The residue measured at 735 °C is lower for washed sisal (11%) than for natural fiber (20%).

In air, this stage corresponds to the combustion of the degraded material. It represents a mass loss of approximately 30% in the natural sisal and 21% in the washed sisal. The $T_{b,COM}$ ranges from 435 to 450 °C, depending on the air flow rate. In the case of washed sisal, it changes from 442 °C, with 50 mL/min air flow, to 432 °C, with a flow of 120 mL/min. The same applies to the exothermic peak temperature, which ranges from 457 °C to 448 °C. Sisal combustion is advanced with respect to that of pure cellulose. Thus, with a 120 mL/min air flow, the $T_{b,COM}$ values of sisal and cellulose are 415 °C and 470 °C, respectively. Comparing the peak temperatures of DTG and DTA signals along the combustion, exothermic effects appear delayed about 12 to 16 °C with respect to the mass loss. A small lag of the heat flow signal in relation to that of mass loss is normal, because the heat is transferred through the crucibles before reaching the thermocouples. In this case, an extra delay is normal because the combustion probably takes place in the upper surface of the sample and thus, the sample itself is an additional barrier for the heat transfer.
Thermal degradation of sisal lignin covers a wide range of temperatures as indicated by the thermal analysis of pure lignin. Since the proportion of this component is very low in the sisal fiber, its decomposition is not defined as clearly as in the case of the other components. The shoulder or tail that appears above 400°C on the DTG curve under inert atmosphere was associated with lignin [52,53]. Mass loss processes with relative maxima at 119, 168, 264, 367 (absolute maximum) and 418 °C were observed for Klason lignin extracted from sisal [33]. In our case, the DTA plot shows a small exothermic effect at around 410 to 435 °C, which is no longer detectable with a high flow of nitrogen.

In the analysis of the thermal decomposition of sisal fiber we must also take into account the analysis of the mineral content. Calcium represents the main component of sisal ashes, as it can be observed in the XRF results in Table 3, whose origin is mainly from calcium oxalate monohydrate, which is accumulated between ultimate fibers of sisal.

Table 3 XRF analysis of Natural and Washed Sisal ashes.

<table>
<thead>
<tr>
<th>% Soluble fraction</th>
<th>Natural sisal</th>
<th>Washed sisal</th>
</tr>
</thead>
<tbody>
<tr>
<td>% Ash</td>
<td>1.2</td>
<td>0.3</td>
</tr>
<tr>
<td>% wt</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CaO</td>
<td>64.60</td>
<td>78.10</td>
</tr>
<tr>
<td>MgO</td>
<td>10.51</td>
<td>14.82</td>
</tr>
<tr>
<td>Na2O</td>
<td>5.92</td>
<td>0.76</td>
</tr>
<tr>
<td>SiO2</td>
<td>4.77</td>
<td>1.56</td>
</tr>
<tr>
<td>Fe2O3</td>
<td>4.12</td>
<td>0.45</td>
</tr>
<tr>
<td>SO3</td>
<td>3.00</td>
<td>1.49</td>
</tr>
<tr>
<td>Al2O3</td>
<td>2.68</td>
<td>0.73</td>
</tr>
<tr>
<td>P2O5</td>
<td>1.85</td>
<td>0.84</td>
</tr>
<tr>
<td>K2O</td>
<td>1.73</td>
<td>0.67</td>
</tr>
<tr>
<td>SrO</td>
<td>0.31</td>
<td>0.24</td>
</tr>
<tr>
<td>TiO2</td>
<td>0.18</td>
<td></td>
</tr>
<tr>
<td>MnO</td>
<td>0.13</td>
<td>0.13</td>
</tr>
<tr>
<td>CaO</td>
<td>0.08</td>
<td>0.08</td>
</tr>
<tr>
<td>BaO</td>
<td>0.06</td>
<td></td>
</tr>
<tr>
<td>ZnO</td>
<td>0.04</td>
<td>0.11</td>
</tr>
<tr>
<td>Cl</td>
<td>0.03</td>
<td>0.02</td>
</tr>
</tbody>
</table>

However, due to the heterogeneity of natural fiber, the small proportion of this inorganic compound, and the superposition of the different degradation stages of natural fiber, not all the decomposition stages of this salt are perceptible,

1) \( \text{CaC}_2\text{O}_4 \cdot \text{H}_2\text{O} \rightarrow \text{CaC}_2\text{O}_4 + \text{H}_2\text{O} \) (near 200 °C)
2) \( \text{CaC}_2\text{O}_4 \rightarrow \text{CaCO}_3 + \text{CO} \) (about 500 °C)
3) \( \text{CaCO}_3 \rightarrow \text{CaO} + \text{CO}_2 \) (close to 800 °C)

The first and third steps are endothermic, independently of the atmosphere. However, the second one is endothermic under inert atmosphere and exothermic in the presence of oxygen, due to immediate oxidation of the generated \( \text{CO} \) to \( \text{CO}_2 \). Thus, just immediately after the combustion stage of sisal in air, the small exotherms accompanied by slight loss of mass in the range from 470 to 480 °C (Table 2), can
be associated to the combined effect of oxalate decomposition and combustion of the resulting CO. It is likely that this effect is also contributed by the combustion of residual carbon originated from fiber lignin. On the other hand, degradation of natural sisal under nitrogen does not include any combustion step and thus, the mass loss associated to the oxalate-carbonate transformation can be clearly observed at 460 °C.

A number of observations can be extracted from above results:

a. The energetic effects match better the mass loss under inert atmosphere than in air. In an oxidizing atmosphere the net exothermic effect is shifted to a higher temperature than its maximum rate of mass loss. The explanation lies in the combination of the various thermal effects caused by the different biomass components, prevailing exothermic volatilization of cellulose, the major component of sisal.
b. The oxidizing atmosphere promotes the pyrolysis stage to lower temperatures.
c. A higher flow rate of purge gas has little effect on the pyrolysis step, which produces a slight advancement of the decomposition temperature in air. In contrast, the increase of the flow rate exerts an important accelerating effect on the combustion stage. Both accelerating effects result from the fact that the presence of oxygen promotes oxidation reactions. The effect of the percentage of oxygen has been studied for the thermal decomposition of diverse lignocellulosic materials [54,55].
d. In comparison with pure cellulose, the temperature of the maximum mass loss rate of the decomposition stage of sisal cellulose, $T_{bCEL}$, is slightly higher. It could be a consequence of an important stabilizing effect of PNC, as well as lignin, which is structurally mixed with cellulose and hemicellulose, as it was deduced for ethanol-extracted woods [56]. On the other hand, the temperature of the combustion stage, $T_{bCOM}$, is lower than that observed for pure cellulose. In an oxidizing atmosphere, combustion gasses (mainly CO$_2$ and H$_2$O) from the lower thermal stability components could facilitate the oxidization of the carbonous residue of the cellulosic fraction of the sisal, thus advancing their combustion process.

3.3.1. Effect of the hot-washing treatment on the thermal decomposition of sisal fiber

From the previous analysis of the thermal behavior of natural and washed sisal, the effect of the washing treatment on the thermal decomposition can be summarized as follows:

a. Hot washing reduces the temperature range in which decomposition occurs. Furthermore, under inert atmosphere, a delay in the pyrolysis of the washed fiber is observed, both at the stage associated with NCP (approximately 13 °C higher in the washed fiber) and cellulose decomposition (between 2 to 4 °C in the washed fiber) (Table 2). These results agree with those found in other studies on washed lignocellulosic material [9,57]. Moreover, washed sisal presents the combustion stage slightly delayed with respect to the natural fiber (Fig. 4b).
b. Washed sisal presents a higher mass loss rate during pyrolysis (Fig. 4a), while a higher combustion rate is observed with natural fiber (Fig. 4b).
c. Decomposition of NCP produces a slightly lower mass loss for washed sisal than for natural fiber, and the opposite occurs due to cellulose decomposition, as is visible in Fig. 4a.
d. No effect of the type of atmosphere on the decomposition of the NCP was observed. However a clear effect was observed in the step associated with the cellulose, and even with a greater intensity, in the case of the washed fiber. The decomposition of sisal cellulose in air is anticipated with respect to pyrolysis under nitrogen, being more notorious in the case of washed fiber: about 20 °C for washed sisal and 12 °C for natural sisal (Table 2).

e. The amount of char generated under inert atmosphere is lower in the washed sisal than in natural sisal (Table 2).

f. The amount of residue obtained in air atmosphere is lower in the case of washed sisal. The washing process produces partial extraction of mineral substances, by partial solubilization. The proportions of Ca and Mg do not change significantly, whereas Ti, Ba, Fe and Na are entirely or almost entirely eliminated by the hot-washing process (Table 3).

Most of the aforementioned effects result from enriching the sample in cellulose as a consequence of the hot-washing removal of substances from the cell walls. The overall result is an increased thermal stability of NCP both in inert atmosphere and in air. The hot-washing process mostly extracts non-cellulosic components of low decomposition temperature, while the higher stability components remain in the fiber, thus increasing the Tb_{NCP} values of the fiber. The slight increase of Tb_{CEL} observed in washed sisal under inert atmosphere would result from a better reorganization and packing of the cellulose crystallites in the fibers, promoted by the removal of certain amount of cementitious material. The crystallinity index of washed sisal (69.7%) and natural sisal (64.6%) confirm that the washing process produces an increase of crystallinity, due to the elimination of cell components. On the other hand, an increase of cellulose thermal stability could also be attributed to the removal of inorganic salts, which could promote its degradation [6]. In contrast, in air, the oxidative pyrolysis hides this effect. The higher percentage of cellulose in washed sisal would explain both the acceleration of mass loss during the second pyrolysis sub-stage, and the delay of the combustion that occurs in an oxidizing atmosphere, getting slightly closer to the behavior of pure cellulose. It is known that the oxidative reactions are responsible for accelerating the depolymerization and mass loss rate of the cellulose decomposition in air [39]. This indicates that in the washed sisal oxidative reactions of cellulose are more favored than in natural fiber, as some protecting components were removed.

The lower percentage of carbon residue generated from washed sisal results from the removal of organic and inorganic matter. The removal of inorganic salts, which were analyzed by XRF, would also indirectly contribute to a lower amount of residue, since they could catalyze the formation of char [58,59].

3.4. FTIR analysis of gases evolved

The three-dimensional diagrams of the infrared absorption spectra of the gases released under inert atmosphere from cellulose, xylan, lignin and hot-washed sisal are shown in Fig. 5. They indicate the evolution of gaseous products as a function of wavenumber and temperature.
Figure 5. Three-dimensional FTIR spectra (Absorbance – wavenumber – Z (temperature, time)) of gases evolved under 120 mL/min nitrogen purge, during decomposition of xylan (a), microcrystalline cellulose (b), lignin (c) and Washed Sisal (d).

The compounds identified in the evolved gases are listed subsequently, specifying the wavenumbers followed: CO$_2$ (2359 cm$^{-1}$), CO (2174 cm$^{-1}$), H$_2$O (3732 cm$^{-1}$), CH$_4$ (3015 cm$^{-1}$), acetic acid (CH$_3$C(O)OH: 1772, 1795, 1273, 1178, 1070 and 1003 cm$^{-1}$), formic acid (H(O)OH: 1795, 1772, 1105 and 1745 cm$^{-1}$), methanol (CH$_3$OH: 1032 cm$^{-1}$), formaldehyde (H(O)H: 2780 [60], 1772, 1745 and 1716 cm$^{-1}$) and some ketone, which tentatively can be assigned to 2-butanone (CH$_3$CH$_2$C(O)CH$_3$: 1745, 1734 and 1178 cm$^{-1}$). Bold type is used to denote the characteristic IR peaks selected for substances monitoring, which were carefully chosen in such form that they correspond to a unique compound, avoiding the overlapping of different gaseous species. In case of coincidences, additional wavenumbers were used to analyze their evolution, which are indicated in italics.

Figure 6 shows the FTIR spectra of gases emitted by washed sisal fiber at two representative temperatures, $T_{bCEL}$ and $T_{bNCP}$, which respectively concern the decomposition of the non-cellulosic
polysaccharides and cellulose. As in the case of microcrystalline cellulose, but to a lesser extent, it is observed a broad band at above 2500 cm\(^{-1}\), which after correction of the spectrum remains at around 3600-3000 cm\(^{-1}\), and a characteristic band at around 1047 cm\(^{-1}\), as indicated by the arrows in Fig. 6. These bands are attributed to a glycol type condensed phase, which was swept by the purge gas, as it will be discussed in the section of gases evolved from cellulose. Furthermore, differently than in the case of the pure biopolymers, a triplet band centered at 858 cm\(^{-1}\) is observed during the decomposition of sisal, denoted by an asterisk in Fig. 6, which is assigned to hydroxyacetaldehyde, (OH)CH\(_2\)C(O)H) [60].

**Figure 6.** FTIR spectra of gases released out during decomposition of Washed Sisal under nitrogen atmosphere, at \(T_b_{PNC}\) (295 °C) and \(T_b_{CEL}\) (350 °C). Arrows denote characteristic bands of the emitted glycol phase, while the asterisk shows the vibration assigned to hydroxyacetaldehyde.

It is known that the absorbance at a specific wavenumber is linearly dependent on the concentration of gas, so the changes in absorbance reflect the tendency of concentration variation of the different gas species, provided that the FTIR procedures are the same [35,42].

Thermograms of Fig. 7, 8, 9 and 10 show the evolution of the main infrared bands associated with gaseous products for pure biomassic components and for washed sisal, which will be discussed in more detail in the following sections. Figure 11 shows the comparison of the evolution of these bands for the different components and the hot-washed sisal, under two flow rates of nitrogen, 50 and 120 mL/min. Data derived from the mentioned thermograms are collected in Tables 4 and 5, which show the relative intensities of different vibrations with respect to the maximum absorbance of CO\(_2\) emitted in the pyrolysis stage for each biocomponent and washed sisal.
Table 4 Maximum intensities of the main infrared bands of volatiles emitted by biomass components, under 120 mL/min of nitrogen or air purges, expressed as relative percentage respect to the maximum absorbance of CO₂ evolved in the respective pyrolysis stages.

<table>
<thead>
<tr>
<th>Cellulose</th>
<th>CO (2174 cm⁻¹)</th>
<th>H₂O (3732 cm⁻¹)</th>
<th>CH₄ (3014 cm⁻¹)</th>
<th>1035 cm⁻¹ Methanol</th>
<th>1105 cm⁻¹ Formaldehyde</th>
<th>2780 cm⁻¹ Acetone</th>
<th>1178 cm⁻¹ Ketone</th>
<th>1734 cm⁻¹ Ketone</th>
<th>1772 cm⁻¹ Acetone</th>
<th>1795 cm⁻¹ Acetone</th>
<th>1047 cm⁻¹ Glycol</th>
</tr>
</thead>
<tbody>
<tr>
<td>N₂</td>
<td>6.0</td>
<td>40.3</td>
<td>4.2</td>
<td>a</td>
<td>55.6</td>
<td>17.8</td>
<td>38.2</td>
<td>107</td>
<td>62.8</td>
<td>47.6</td>
<td>105</td>
</tr>
<tr>
<td>Air</td>
<td>4.8</td>
<td>13.1</td>
<td>N.D.</td>
<td>a</td>
<td>5.7</td>
<td>2.7</td>
<td>2.1</td>
<td>22.8</td>
<td>16.5</td>
<td>16.2</td>
<td>9.2</td>
</tr>
<tr>
<td>Xylan</td>
<td>N₂</td>
<td>11.5</td>
<td>10.2</td>
<td>5.6</td>
<td>9.3</td>
<td>13.8</td>
<td>0.9</td>
<td>4.2/6.4</td>
<td>33.4</td>
<td>9.9/26.9</td>
<td>9.9/23.9</td>
</tr>
<tr>
<td>Air</td>
<td>5.7</td>
<td>13.4</td>
<td>N.D.</td>
<td>9.1</td>
<td>13.7</td>
<td>1.6</td>
<td>1.9/6.9</td>
<td>26.3</td>
<td>3.2/24.0</td>
<td>3.5/20.5</td>
<td>N.D.</td>
</tr>
<tr>
<td>Lignin</td>
<td>N₂</td>
<td>2.2</td>
<td>15.4</td>
<td>11.7/18.8</td>
<td>14.0</td>
<td>N.D.</td>
<td>N.D.</td>
<td>d</td>
<td>d</td>
<td>d</td>
<td>N.D.</td>
</tr>
</tbody>
</table>

/ Two stages can be distinguished
N.D. Vibration not detected
a If vibration exists is hidden by the glycol band
b Its intensity is increased due to the glycol band
c Acetic acid contribution doesn’t exist
d Only water vibrations are observable

Table 5 Maximum intensities of the main infrared bands of volatiles emitted by Washed Sisal, under different atmosphere and flow rate, expressed as relative percentage respect to the maximum absorbance of CO₂ evolved in the respective pyrolysis stages.

<table>
<thead>
<tr>
<th>Washed Sisal</th>
<th>CO₂ (2359 cm⁻¹)</th>
<th>CO (2174 cm⁻¹)</th>
<th>H₂O (3732 cm⁻¹)</th>
<th>CH₄ (3014 cm⁻¹)</th>
<th>1035 cm⁻¹ Methanol</th>
<th>1105 cm⁻¹ Formaldehyde</th>
<th>2780 cm⁻¹ Acetone</th>
<th>1178 cm⁻¹ Ketone</th>
<th>1734 cm⁻¹ Ketone</th>
<th>1772 cm⁻¹ Acetone</th>
<th>1795 cm⁻¹ Acetone</th>
<th>1047 cm⁻¹ Glycol</th>
</tr>
</thead>
<tbody>
<tr>
<td>N₂ 50 mL/min</td>
<td>420.0/100</td>
<td>1.8/8.9</td>
<td>7.7/19.1</td>
<td>5.1/5.3</td>
<td>11.4/22.2</td>
<td>11.3/39.0</td>
<td>14.0/2.4</td>
<td>848/48.6</td>
<td>270/94.3</td>
<td>139/111</td>
<td>160/81.5</td>
<td>9.5/36.3</td>
</tr>
<tr>
<td>Air 50 mL/min</td>
<td>18.8/100</td>
<td>0.9/4.5</td>
<td>1.7/9.8</td>
<td>N.D.</td>
<td>2.8/36</td>
<td>2.7/54</td>
<td>N.D.</td>
<td>17.7/11.3</td>
<td>1.3/12.6</td>
<td>27.6/24.3</td>
<td>33.4/24.0</td>
<td>2.3/4.6</td>
</tr>
<tr>
<td>N₂ 120 mL/min</td>
<td>40.0/100</td>
<td>1.6/8.4</td>
<td>7.9/17.1</td>
<td>5.5/5.9</td>
<td>10.7/32.5</td>
<td>13.9/56.6</td>
<td>1.3/1.3</td>
<td>92.2/54.5</td>
<td>29.5/117</td>
<td>138/121</td>
<td>160/79.3</td>
<td>9.1/50.8</td>
</tr>
<tr>
<td>Air 120 mL/min</td>
<td>26.5/100</td>
<td>0.7/3.9</td>
<td>3.6/14.2</td>
<td>N.D.</td>
<td>1.9/40</td>
<td>2.3/67</td>
<td>0.2/2.1</td>
<td>15.4/9.5</td>
<td>7.0/28.9</td>
<td>24.3/30.0</td>
<td>27.0/23.3</td>
<td>1.6/6.0</td>
</tr>
</tbody>
</table>

/ NCP and cellulose stages are distinguished
ND Vibration is not detected
a Its intensity is increased due to the glycol band
b Overlapping with Acetic acid band
3.4.1. Analysis of gases evolved from the pure lignocellulosic components

The analysis of gases evolved in the decomposition of pure components in air and nitrogen is described, previously to study the more complex case of sisal fiber.

i) Xylan:

In nitrogen atmosphere, the following species are detected at temperatures below 350°C: CO₂, CO, H₂O, acetic acid, formic acid, methanol, formaldehyde and a ketone, with vibrations near 1178 and 1734 cm⁻¹ (Fig. 7).

Figure 7. Evolution of the main infrared bands of the volatiles emitted during xylan decomposition, under 120 mL/min nitrogen (a-c) and air (d-f) purges, with the corresponding DTG curve.

Among the three biomass components considered, xylan is the one which presents the highest absorbance of CO₂ and the highest CO/CO₂ ratio (Table 4). CO₂ comes from the breakdown of C=O and COOH groups [42]. It would mainly come from decarboxylation reactions of O-acetyl units according to
Shen et al. [61]. In the present case that contribution must be less important, since the starting material is a hemicellulose mostly composed of xylose units. The CO formation is attributed to decarbonylation reactions, specifically to the breakdown of COC and C=O [42], due to the decomposition of the ring-opened intermediate products [61]. The H2O emission during the pyrolysis stage is mainly derived from the hydroxyl groups of xylopyranose units. The emission of acetic acid and methanol is attributed to the breakdown of the O-acetyl and O-methyl groups of the residual side chains, which are even present after the extraction and purification of commercial xylan. The presence of various ketones in the products resulting from xylan pyrolysis was identified using Py-GC/MS by Y. Wu et al. [62], who found at 260 °C mainly 1-hydroxy-2-butanol, 1-hydroxy-propanone, and, to a lesser extent, 4-hydroxy-3-hexanone.

In nitrogen at temperatures above 400 °C certain gases generated by secondary reactions of decomposition of anhydrous sugars are distinguished. Thus, the CO2 emission continues, but slightly, and CH4 is detected with a maximum near 470 °C, close to the exothermic effect observed in the DTA of xylan (Table 1). CH4 is formed by breaking R-CH3 bonds, which requires elevated temperatures due to its high stability [61]. The CO2 emission starts to increase at 550 °C, reaching a maximum rate at about 700 °C, temperature at which a new CO emission begins.

During the pyrolytic decomposition stage in air the same gases are detected than when using inert atmosphere. Their proportions with respect to the CO2 emitted are quite similar to those obtained under nitrogen atmosphere, with the exception of CO principally, whose emission diminishes in air (Table 4). Furthermore, mass losses in air and nitrogen are similar, 59 and 63% respectively (Table 1). It can be explained through the phenomena that affect the volatiles emitted in the decomposition step. In this case, the decomposition temperature of xylan (about 290 °C) is sufficiently low so that, firstly, most of the volatiles generated are not oxidized to CO or CO2 in air, and on the other hand, the self-generated CO would oxidize to CO2 in contact with air through the thermodynamically favored reaction: 2CO + O2 \rightarrow 2CO2, which explains the lower proportion of CO. The CO2 emission during the combustion process, between 300 – 420 °C, is intense and presents an asymmetrical shape which indicates that the process is triggered by the temperature once the energy barrier of the combustion process is reached. Then, the combustion rate decreases as the remaining amount of sample decreases. The corresponding DTG plot reflects the same process, but the DTG peak is more resolved due to the delay and mixing of components that may occur during the gas flow from the TGA furnace to the FTIR detector [63]. Additionally, small emissions of CO2, H2O and CO, associated with mild exothermic effects are observed at around 440 °C and at 570 °C (Fig. 7D and Table 4). The exotherm at 440 °C corresponds to the completion of the 300 – 420 °C combustion process, which produced an overheating on the programmed ramp. Then, the process was halted as the temperature decreased to recover the programmed values, as it can be observed on the DTG plot. The exotherm at 570 °C corresponds to combustion of a more stable carbon residue, which resulted from degradation of the most stable structures.

**ii) Cellulose:**

In nitrogen, at temperatures below 400 °C, CO2, CO, H2O, formaldehyde, formic acid, and 2-butanone are detected, besides a glycol condensate (Fig. 8). This condensate is assigned to 1,2-
propanediol, intermediate reaction product, and to ethylene glycol, which was observed in pyrolytic decomposition of cellulose [64,65]. It should be noted that the signal of formic acid (1105 cm\(^{-1}\)) is increased due to the glycol (1047 cm\(^{-1}\), Fig. 8B and 11C), which also hides methanol (1032 cm\(^{-1}\)). Thus, it is not possible to determine the proportion of each component (Fig. 8A and 11F, and Table 4).

Figure 8. Evolution of the main infrared bands of the volatiles emitted during microcrystalline cellulose decomposition, under 120mL/min nitrogen (a-c) and air (d-f) purges, with the corresponding DTG curve.

In contrast to xylan most of the main gaseous products of cellulose are evolved together, in a narrower range of temperatures, with maximum close to 330 - 340 °C, except for CH\(_4\), which is emitted at higher temperature. Cellulose is the component that produces the highest emissions of H\(_2\)O and organic products with respect to the CO\(_2\) emitted (Table 4). The breaking of the glycosidic linkages and ring opening generate liquid and gaseous intermediates, whose identity and proportion are dependent on the pyrolysis conditions. Shafizadeh [66] and subsequently Shen et al. [67] collected and proposed different routes of formation of CO\(_2\), CO, CH\(_4\), formaldehyde, methanol and acetic acid, among other products of
decomposition of cellulose. In our study acetic acid has not been detected, so the bands located at 1178, 1772 and 1795 cm⁻¹, shown in Fig. 8B, 8C, 11G, 11H and 11L are associated to acetone, formaldehyde and formic acid. Shen et al. [67] indicate that the formation of CO is favored by high temperatures and long residence times, which explain that CO formation is mainly affected by the secondary reactions, especially of aldehyde-type low molecular weight compounds. Nevertheless, CO₂ is mainly formed in the early stages of the pyrolysis of cellulose. Li et al. [60] found that formaldehyde formation is favored by a lower purge flow and by longer residence times, which would explain its formation through secondary reactions. They explain that at high temperatures, near 550 °C, formaldehyde decomposes to generate CO and H₂, which is consistent with the observation of Shen et al. Various ketones have been proposed and detected at the primary decomposition and pyrolysis of cellulose. Thus, 2,3-butanedione is among the products of pyrolysis of levoglucosan, a major decomposition product of cellulose [68]. 2-butanone, detected here at temperatures below 400 °C, has also been detected by Py-GC/MS analysis [62]. According to other reports, it may come from decarboxylation of levulinic acid [66]. Radlein et al. [69] found formic acid in the pyrolysis products of different celluloses, which would be originated from the decomposition of levoglucosan according to Hosoya et al. [70].

In nitrogen, at temperatures above 450 °C, CO₂, CO and CH₄ are detected, resulting from side reactions. In this stage, maximum emission of CO₂ and CH₄ is observed at about 530 °C, while the maxima of CO₂ and CO appear at 650°C, as shown in Fig. 8A, 11A, 11E and 11I.

The products detected during the decomposition in air of cellulose are the same than under inert atmosphere. However, compared to that of CO₂, the emission of organic compounds and H₂O has decreased significantly in air, while the proportion of CO is maintained in almost the same proportion as in nitrogen (Table 4). Since the mass loss after pyrolysis under air and nitrogen are similar, the explanation lies primarily in the changes suffered by the emitted gaseous compound.

In an oxidizing atmosphere, when the temperature is high enough, it could happen that at the same instant in which the decomposition/pyrolysis of the cellulose occurs, the volatile organics generated oxidize, which would be favored by the presence of steam. Thus, a gasification reaction explains the lower proportion of H₂O detected. On the other hand, the consumption of CO according to the reaction 2CO + O₂ → 2CO₂ is less thermodynamically favored, which explains its higher relative proportion under air atmosphere. The evolution of CO₂ in the combustion process comprises a wide interval of temperatures, from 370 to 700 °C and maximum at 470 °C, where CO and H₂O are detected. Exothermic effects, around 540 and 600 °C, detected in the DTA of the cellulose, correspond mainly to the emission of CO₂, as observed in Fig. 8B, associated to additional combustion phenomena.

iii) Lignin:

Unlike cellulose and hemicellulose, degradation of lignin under inert atmosphere covers a very wide range of temperature, from 130 to 500 °C (Fig. 9). The gases detected in this interval are CO₂, CO, H₂O, CH₄ and CH₃OH. No other organic compounds were detected.
Of the three biomass components, lignin is the one which produces the highest proportion of CH₄ and CH₃OH and the lowest of CO with respect to the maximum absorbance of CO₂ emitted at the pyrolysis stage (Table 4). At temperatures below 500 °C, the emission of H₂O can be associated to aliphatic and aromatic hydroxyl groups, and the emission of CO₂ and CO to cracking and reforming of the carboxyl, carbonyl and ester groups, present in the side chains of the phenylpropane units [41]. Their evolution above 500 °C is associated with the secondary pyrolysis of volatiles. In particular, several authors claim that CO is originated at the breaking of two different types of ether bonds: at low temperature the ether bonds between different lignin subunits, whilst at high temperature diaryl ether groups are involved [71,72]. Below 400°C, the evolution of CH₃OH and CH₄ is mainly associated with the cleavage and demethylation of methoxyl groups (-O-CH₃) present in lignin [42,73]. Methanol
formation may also be contributed by the -CH₂OH groups located on the γ carbon of the alkyl lateral chain of the phenylpropane units [41,71]. Above 400°C methane emission continues, reaching a second maximum at 525 °C, more intense than the first one. It is produced by side reactions, and also by primary reactions of condensation and reorganization of the char. At these high temperatures the rupture of the aromatic rings starts, generating methyl radicals and methylene groups, and finally forming methane [41] [71]. From 600 °C methane signal begins to disappear, because it breaks down into H₂ and C [72].

The composition and evolution of gases emitted by lignin in this study turn out to be virtually identical to those of alkaline lignin studied by Yang et al. [42]. Other studies describe the emission of other organic compounds, not detected in the present work, such as formaldehyde, formic acid and phenols originated by the rupture of phenylpropane units. The type of lignin and the extraction method would explain the differences in the composition of the emitted gases. The basic reason why a smaller variety of volatile organic compounds is detected is that char formation reactions are favored against the generation of volatile compounds. G type lignins, as the one used in this study are prone to undergo condensation and coupling reactions between phenylpropane units, causing them to generate more amount of char and emit less amount of phenolic derivatives [41], which would be characterized by intense vibrations around 1500, 1250 and 1100 cm⁻¹ [73]. On the other hand, lignins containing little sulfur, as the one studied, are more affected by this type of condensation reactions. Fenner and Lephardt [73] indicate that the incorporation of sulfur facilitates the fragmentation of phenylpropane units and decreases its condensation.

3.4.2. Analysis of the gases emitted by hot-washed sisal fiber

The evolution with temperature of the main infrared bands of the gases emitted during the decomposition under inert and oxidant atmosphere of washed sisal fibers is shown in Fig. 10. The gaseous components observed at different stages are summarized below:

i) At temperatures below 150 °C, H₂O emission is mainly from to the moisture in the fibers.

ii) In the first sub-step, below 300 °C, the gases CO, CO₂ and H₂O come from the decomposition of sisal lignin and mainly from hemicelluloses and other NCP still present in the washed sisal. The emission of acetic acid, formic acid, methanol, formaldehyde, and some ketone, which may be tentatively assigned to 2-butanone, is mainly due to decomposition of hemicellulose. Sisal lignin should also contribute to the emission of methanol and acetic acid as suggested by other studies with HGS type lignins [41]. Nevertheless, in studies with wood, the emission of formic acid at this stage has been associated only with the hemicellulose and not with lignin [74]. Notably, the proportion of acetic acid emitted (Table 5) is proportionally much higher than that observed in the pure xylan (Table 4). This is because sisal hemicelluloses contain more side chains with acetyl groups than the xylan studied.
In the second sub-stage, between 300 and 400 °C, the emitted compounds are mainly due to the pyrolysis of sisal cellulose: CO, CO₂, H₂O, glycol, formaldehyde, methanol, formic acid and ketone, assigned tentatively to 2-butanone, as well as hydroxyacetaldehyde which starts to be generated in the previous stage. The latter compound may appear as a product of decomposition of cellulose by a competitive route to the formation of levoglucosan, the main component of cellulose tar [60,67]. The low amount of hydroxyacetaldehyde emitted during the pyrolysis of sisal cellulose suggests a route of decomposition different from that of the microcrystalline cellulose studied in this work, in which this compound was not observed. The maximum emissions of all these compounds match with the maximum mass loss rate, as shown in Fig. 10. Acetic acid bands decrease very quickly at the end of the first sub-stage. Thus, the vibrations of the carbonyl groups that remain in the second sub-step can be only assigned to formaldehyde, formic and 2-butanone (Fig. 10C). To a lesser extent sisal lignin contribute to the
emission of CO, CO$_2$ and H$_2$O, and probably also to methanol, formic acid and acetic acid. The emission of CH$_4$ in nitrogen starts at around 360°C (Fig. 10A and 11I). It is mainly associated with the rupture of the methoxyl groups of lignin, but its signal is weak, being obscured by the C-H stretching vibrations of other organic compounds. Additionally, at the end of this sub-stage in air, the combustion of sisal hemicellulose and other NCP occurs, emitting CO$_2$ and H$_2$O (Fig. 10D).

Figure 11. Comparison of the evolution of the main infrared bands of gaseous compounds evolved under nitrogen atmosphere, during decomposition of biomass components under 120 mL/min purges, and decomposition of Washed Sisal under purges of 50 and 120 mL/min. The asterisk of Fig. 11i pyrogram on the first stage of CH$_4$ evolution corresponds to background signal and not to methane vibration.

iii) Both in nitrogen and in air, emission of most organic compounds disappears at temperatures above 400 °C. Under inert atmosphere, the evolution of gases is similar to that of pure cellulose, as it is the majority component. Emission of CO$_2$ and CO start again at 400 °C due to side reactions, being the growth of CO intensity much slower. Emission of CH$_4$ is also observed to increase at 400 °C, whose signal remains at a nearly constant value up to 580 °C, when it began to decrease and a new increase of
CO is observed. The methane signal is zero due to its transformation into H₂ and C, at about 700 °C for a nitrogen flow of 50 mL/min, and 650 °C for 120 mL/min of N₂ (Fig. 11I). The CO₂ and CO reach maximum intensity at around 660 °C with a flow of 50 mL/min, and around 630 and 610°C with a flow of 120 mL / min (Fig. 11A and 11E). These temperatures nearly match the DTA exotherm peaks at 650 and 635 °C for 50 mL/min and 120 mL/min, respectively (Table 2).

In air, combustion occurs and maximum CO₂ emission, accompanied by H₂O and CO, is observed at the maximum rate of mass loss, at around 440 and 430 °C for 50 and 120 mL/min flow rate, respectively (Fig. 10D). The emission of CO₂ continues to 500 - 550 °C, presenting a small relative maximum in the 460 to 470 °C range. This relative maximum matches with mild exothermic processes (around 480 °C) observed on the DTA curve, which have been associated with the decomposition of calcium oxalate and the oxidation of CO to CO₂, as already explained. Also, it is very likely that the flue gas of residual carbon generated by the lignin contribute to these signals.

In light of the results obtained the following remarks can be made:

a. The gaseous products emitted at temperatures below 400 °C for the decomposition of sisal, are basically the same in both oxidizing and inert atmospheres, but in different proportions. During oxidative pyrolysis, these compounds are emitted to a lesser extent with respect to the CO₂ emitted in this stage (Table 5). The same behavior was observed in the case of pure cellulose (Table 4). However, the mass losses observed in each one of the two decomposition sub-stages (NCP and cellulose) are similar independently of the nature and flow rate of the purge considered: 17 and 60% for 50 mL/min of air, 17.6 and 58.5% for 120 mL/min of air, 18.5 and 63.6% for 50 mL/min of N₂, 17.8 and 59.4% for 120 mL/min of N₂ (Table 2). The effect of the oxygen proportion is perceptible in gas phase. Decomposition temperatures for both sub-stages are enough for the generated gaseous products to oxidize immediately, generating CO₂, favored by the presence of steam. Thus, a higher emission of CO₂ is observed under air atmosphere. With nitrogen purge, although to a lesser extent, the oxidation of volatiles is also possible. The reason is that air goes into the furnace when inserting the sample. Although a purge of nitrogen sweeps off most of the air, a little amount of oxygen may always remain in the furnace. Thus, a higher flow of inert gas is more efficient in preventing the oxidation of volatiles and also of the solid residue.

The presence of constituents as cellulose, lignin and mineral salts, would explain the different proportions of organic compounds emitted during the first sub-stage of sisal decomposition according to the kind of atmosphere used, fact not observed in pure xylan. Furthermore, the increase in the decomposition temperature of the hemicellulose could be enough to elevate the activation threshold to initiate the oxidization of volatile compounds.

b. An increased flow of purge gas does not produce any change in the nature of the flue gases. However, it exerts a number of effects in the evolution of gases, which are summarized below:

b.1. Dilution of the output gas. The FTIR spectra show, in general, that the absolute absorbance values are slightly lower in the spectra obtained at higher flow (Table 5 and Fig. 11). A greater flow rate decreases the concentration of compounds emitted.
b.2. Decreasing the intensity of the secondary processes experienced by the volatile products, by shortening the residence times in the furnace. For example, it can be observed in Fig. 11A how a higher flow rate reduces the oxidation of the volatiles at temperatures above 450 ºC, and thus reducing the CO₂ formation.

b.3. Advancement of the apparent temperature range in which the gases generated by side reactions are emitted (e.g., the CO emission above 450°C, Fig. 11E).

b.4. As disadvantage, a higher flow carries a higher percentage of glycol condensate, increasing the infrared vibration intensity by overlapping, like methanol and formic acid signals and, even, hiding them, as occur in the case of volatiles emitted by pure cellulose.

4. Conclusions

TG-FTIR analysis of pure biomass components has helped to clarify the thermal decomposition phenomena of sisal fibers, both in nitrogen and air atmospheres. This analysis may be extended to other lignocellulosic fibers.

Thermal behavior of sisal is explained as a combination of its main biomass components, prevailing cellulose. The research confirms that other constituents, such as non-cellulosic polysaccharides (NCP) and mineral matter, modify the thermal behavior of the main component. The presence of pectins, hemicelluloses and inorganic elements exerts different effects on the temperature of pyrolysis and combustion of sisal cellulose. On one hand, the protective effect of the NCP explains the delay in the pyrolysis stage of it. However, in an oxidizing atmosphere, the flue gases from the NCP facilitate the oxidation of the char, thus advancing the combustion process. The hydroxyacetalddehyde detected during the pyrolysis stage of sisal cellulose, and not in pure cellulose, indicates that the interaction with other components enables cellulose decomposition through an alternative route to that of levoglucosan generation.

The hot washing reduces the range of temperature in which sisal decomposition occurs. An increased thermal stability of sisal NCP, both under inert and air atmosphere, is produced by the hot washing process due to the removing of substances from the cell walls. Furthermore, enriching the sample in cellulose produces an increase of the mass loss rate at the second sub-step of pyrolysis, a delay of the combustion process and, depending on the decomposition atmosphere, different temperature displacements. Thus, under inert atmosphere, a slight increase of the temperature at the maximum mass loss rate of cellulose decomposition (TbCEL) is observed, due to a better packing of the cellulose crystallites and removal of inorganic salts. Under an oxidizing atmosphere, this effect is not observed and TbCEL is shifted to lower temperatures, as cellulose depolymerization is accelerated by oxidative reactions favored in washed sisal once the protective components are removed.

Increasing the air flow produces a slight advancement of the oxidative pyrolysis and a significant advancement of combustion. Respect to the gas evolution, an increased flow results in dilution of the exhaust gases, decreasing the FTIR absorbance signals, besides reducing the intensity of the reactions of volatile products in the furnace, and advancing the apparent temperature range at which the resulting gases are emitted.
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