STUDY BY INFRARED SPECTROSCOPY 
AND THERMOGRAVIMETRIC ANALYSIS 
OF TANNINS AND TANNIC ACID

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ABSTRACT

Among the compounds in nature useful for people are tannins, which present large molecular complexity and they had have an ancient use as tanning agents, and nowadays some of them are used due their antioxidant properties for protect metals of corrosion. Also elucidation, synthesis and purification represent a great challenge to obtain them. Due this fact researches have tried to copy the synthesis of less complex compounds similar to tannins. From a different point of view, the use of tannins extracted from trees bark, considered in many places as a waste, grant an extra value to the bark. In this work we have studied and analyzed through gel permeation chromatography (GPC), infrared spectroscopy (FT-IR) and thermogravimetric analysis (TGA) the average molecular weight, main functional groups and thermal properties of tannins extracted from Schinopsis quebracho colorado. The results of these analyses are compared with those obtained for the commercial grade tannic acid. The comparison allowed to corroborate that is possible to use tannins from the Schinopsis quebracho colorado bark, with functional groups and thermal properties which tend to be similar to the tannic acid. www.relaquim.com

Keywords: coating, corrosion, Schinopsis quebracho colorado, tannic acid, tannins.

RESUMEN

Entre los compuestos que existen en la naturaleza útil para la población, se encuentran los taninos, los cuales presentan una gran complejidad molecular y han tenido un uso desde la antigüedad como agentes para curtido. Hoy en día algunos de ellos se utilizan por sus propiedades antioxidantes para proteger los metales de la corrosión. Además la elucidación, síntesis y purificación representa un gran

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INTRODUCTION

Tannic acid and tannins are polyphenolic compounds able to build very complex chemical structures. The tannic acid, which has an estimated chemical formula C_{76}H_{52}O_{46}, is a commercial form of tannins. 300 years ago Egyptians extracted these compounds and used them as tanning agents. Due their tanning capacity these compounds were very important at an industrial level and tanneries. Although nowadays salts and synthetic tannins have acquired more importance against tannins extracted from natural sources (Matamala et al., 1994; Haslam, 1996).

In the last two decades the isolation and structural determination of many types of tannins, which also include hydrolysable and condensable tannins have been possible using techniques as UV-visible, nuclear magnetic resonance (¹H-NMR, ¹³C-NMR), mass spectroscopy (Isaza et al., 2004; Meagher et al., 2004) and soon. According to their structural differences between many species of tannins, some researchers have been used these compounds for different applications within which are find as fungicide, antibiotic and antioxidant (Schofield et al., 2001; Latté and Kolodziej, 2004). The antioxidant property of tannins has awaked special interest in the corrosion area, due to the fact that these compounds can inhibit corrosion in metals when they are incorporate in coatings. It can find in literature many researches about corrosion inhibition using tannic acid or tannins extracted i.e. from pine, mimosa and acacia species (Jaén et al., 2003; Hernes et al., 2004). The antioxidant effect of these polyphenols compounds is attributed at their quelating action over transition metals and their capacity to attach free radicals through the synthesis of less reactive structures like show Iglesias et al. (2001). It is worth to mention that tannins extraction from trees bark offer an extra value to the barks, which is considered in many places as a waste. Even when tannins formula varies among tree species, tannins present structures similar than the tannic acid, for this reason elucidation and comparison between these compounds is important.

The aim of this work is to corroborate by gel permeation chromatography, infrared spectroscopy and thermogravimetric analysis that tannins extracted industrially (tannic acid), and tannins extracted from Schinopsis quebracho colorado have almost thermal, chemical and physicochemical si-
milar characteristics, and therefore tannins can be reliable compounds for used i.e in anticorrosive coatings.

**MATERIALS AND METHODS**

In this study it has used tannic acid supplied by Sigma-Aldrich. Condensed tannins (98%) were provided by FITECMA-UMSNH (Facultad de Ingenieria en Tecnologia de la Madera de la Universidad Michoacana de San Nicolás de Hidalgo) and they were extracted from *Schinopsis quebracho colorado* bark based in the works of Isaza et al. (2004) and Sivakumaran et al. (2006). Tannins were milled to generate size smaller particles (100 micrometers).

The molecular weight of tannins was obtained by high pressure liquid chromatography (HPLC) with a gel permeation chromatography system, 717 Plus Autosample model using dimethylformamide as solvent. Structural characteristics of the compounds were analyzed with a FT-IR spectrometer (Nicolet 5ZDA) in the absorption mode, ranging from 4000 to 400 cm\(^{-1}\). Thermogravimetric analyses were performed at a heating rate of 10\(^{\circ}\)/min under nitrogen flow by a differential scanning calorimetric (DSC) 822 Toledo Mettler.

**RESULT AND DISCUSSION**

The number average molecular weights (Mw) obtained through GPC for tannins from *Schinopsis quebracho colorado* and tannic acid (Sigma-Aldrich) were of 6 485 and 1 701.23 respectively. The molecular weight of tannins is approximately four times higher than acid tannic because a lot of hydroxyl benzene rings are connected with similar structures forming a big structure as is suggested by Grassino et al. (1999) and Meagher et al. (2004).

The Figure1 shows the spectrum of tannins and tannic acid where it can find a strong absorption around 3700 and 3000 cm\(^{-1}\) with a wide and strong band centered at 3381 and 3356 cm\(^{-1}\) respectively. These bands are assigned to the hydroxyl groups (OH) stretching vibrations and due to the wide variety of hydrogen bonding between OH.

In this spectrum it can notice a sharp peak at 2927 and a small shoulder at 2860 cm\(^{-1}\) associated with the symmetric and antisymmetric \(-C-H-\) stretching vibrations of CH\(_2\) and CH\(_3\) groups respectively. Also it does not appear signals of free water molecules due there is no evidence of bands at 2130 cm\(^{-1}\) and 1650 cm\(^{-1}\) (H-O-H bending vibrations) and at about 700 cm\(^{-1}\); but in

![Figure1. Infrared spectroscopy of tannins and tannic acid](image-url)
the tannic acid spectrum some of these peaks appear due some hydrogen and oxygen atoms in this molecule can be bonding and they are observed in the spectrum as water signals.

The deformation vibration of the carbon-carbon bonds in the phenolic groups absorbs in the region of 1500-1400 cm⁻¹. At 754 cm⁻¹ shows the result distortion vibration of C=C in benzene rings (Socrates, 2004). Around 1452 cm⁻¹ stretching vibrations of -C-C aromatic groups appear in both spectrums respectively.

Spectrum of tannins compared with tannic acid and tannins shows a decrease in many peaks located at 1718, 1614, 1452, 1328 and 1191 cm⁻¹ which are not sharp and strong as is observed in tannic acid. Also it can be observed that only the tannic acid contains some aromatic esters due to the signal characteristics bands of carbonyl groups: C=O stretching vibration at 1730-1705 cm⁻¹ and C-O at 1100-1300 cm⁻¹ (Silverstein et al., 1981; Stuart, 2005).

Detailed information is listed in Table 1; these results are similar to those reported in the literature for polyphenolic compounds as tannic acid (Socrates, 2004).

The TGA traces and mass spectra obtained for the tannins and tannic acid heated at a rate of 5 °C cm⁻¹ are shown in the Figures 2(a)-(b), which show the dependence of the mass loss of the samples expressed as a percentage of the initial mass and temperature. Also the first derivative is below of them.

Two distinct mass loss peaks can be seen in the Fig. 2(a): a weak peak centered at 48.8 °C where almost 11.47% of weight due the postcuring, thermal reforming, preliminary oxidation steps and elimination of volatile fractions. The second peak, is sharper and more pronounced and it is found at 249.9 °C where the degradation tannins begins and it could be the result of the partial breakdown of the intermolecular bonding. Finally after at 500 °C remains a carbonized residue of 55.29% w.

TGA of tannic acid in Fig. 2 (b) shows five thermal zones, the first one is appreciated between 25 and 150°C with a mass loss centered at 37.7 °C where preliminary oxidation steps and elimination of volatile fractions occurs.

In the second zone (151 and 205 °C) is carried out the first depolymerization of tannic acid approximately 1.004%w. Between 206 and 295 °C is presented a second

Table 1. Infrared positions of tannins and tannic acid.

<table>
<thead>
<tr>
<th>Group</th>
<th>Vibration</th>
<th>Approach frequency/experimental cm⁻¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arom-CH₂-OH</td>
<td>v OH</td>
<td>3400/3381</td>
</tr>
<tr>
<td>Arom-CH₂-OH</td>
<td>v₂ CH₂</td>
<td>2920/2925</td>
</tr>
<tr>
<td>Arom-C-O-</td>
<td>v C-O</td>
<td>1630/1614</td>
</tr>
<tr>
<td>Arom-CH₂-OH</td>
<td>δ CH₂</td>
<td>1420/1452</td>
</tr>
<tr>
<td>Arom-O-</td>
<td>v₁ C-O</td>
<td>1050/1165</td>
</tr>
<tr>
<td>Arom-O-</td>
<td>v C-O</td>
<td>800/646</td>
</tr>
</tbody>
</table>

*Socrates, 2004
depolymerization of the material (36.62%w) with a pronounced peak at 255.49°C. Third degradation of tannic acid takes place after 296 °C with a remarked peak at 306.39 °C; in this section is seen a mass loss of 26.41%w. Finally after 500 °C remains 28.3% w of carbon.

Comparing differences between tannic acid and tannins, we observe in the last one, a difference of more than 27% weight of carbon residues, due tannins contain impurity traces. Also, tannins contain more atoms of carbon in their structure than tannic acid. Another important difference between tannins and tannic acid thermograms, is that in the first one we observe three thermal zones while in tannic acid it observe five zones, due tannic acid has a less complex structure, and its bonds can be easily broken forming smaller structures.

CONCLUSIONS

In this work we have studied the molecular weight, main functional groups and thermal stability of a natural product, tannins from Schinopsis quebracho colorado and it was compared with results of a similar synthetic compound (tannic acid, Aldrich®). Trough infrared spectroscopy it was observed that tannins do not shows important signal from carbonyl groups (aromatic esters) while they appears in tannic acid. Another important difference is in the hydroxyl group band, which in tannins is less pronounced, because several hydroxyls are interacting among them.

Thermogravimetric analysis shows that tannins show a similar stability thermal that tannic acid, which also have more depolymerization stages due to the easily break in smaller structures than tannins (compound con some monomer units similar to the tannic acid).

Even when results of GPC, FT-IR and TGA vary slightly, it can be observed that Schinopsis quebracho colorado tannins present structures and functional groups which tend to be similar to the tannic acid. The above makes tannins ideal candidates to give them a good application, for example as anticorrosive coatings.

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