CHITOSAN METAL COMPLEXES AND CHITOSAN-Cu ESR STUDIES

EDELIO TABOADA VALDÉS¹, GALO CARDENAS TRIVIÑO

¹Escuela de Ingeniería Ambiental, Facultad de Ingeniería, Universidad Católica de Temuco, Temuco, Chile, etaboada@uct.cl
Galo Cárdenas Triviño*, ²Laboratorio de Materiales Avanzados, Departamento de Polímeros Universidad de Concepción, Casilla 160-C, Concepción, Chile.
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ABSTRACT

Properties of the polymer metal complexes can be explained up to certain extent by the molecular structure of the complexes, but this area has not been well studied in the case of the chitosan metal complexes. In this work, we are proposing that chitosan is bonded through the 4 nitrogen to copper by means of square planar geometry. The polymer requires some structural modifications in order to be arranged in that geometry. Two sites in the chitin were detected, one similar to chitosan and the other in a square planar tetrahedrally distorted arrangement, with three atoms of nitrogen and one of oxygen or two atoms of nitrogen and two of oxygen. ESR studies of the copper complexes by means of chlorine and nitrate as ion counter were carried out. Nitrate exhibits one type of retention site and, on the other hand, chlorine presents two types of retention sites.

Keywords: Chitosan, metal complexes, esr, retention sites.

INTRODUCTION

Several publications on chitosan complexes have been carried out in order to trap metal ions at different pH³. Chitosan phosphate and chitosan dithiocarbamate were synthesized by Lopez de Alba ⁴ to trap uranium (VI). Lasko ⁵ obtained derivatives from mercaptosuccinic acid, pyridoxal hydrochloride and succinamide and a comparison with chitosan retention against Cu (II), Cd (II), Fe(II) and Pb(II) was carried out. Guibal has prepared a graft copolymer with oxo-2-glutaric acid in order to trap both uranium and vanadium ⁶.

Among the few attempts to understand more about the structure that different metal ions produces with chitosan, it must be noted the work reported by Schlick ⁷, in which an ESR study on the Cu (II) bond sites in chitin and chitosan, respectively was performed.

Other similar studies using this technique, combined with Mössbauer spectroscopy, proposed a square planar structure for Cu(II) ion and a distorted rombic array for the Fe(III) complex with chitosan⁸. Mössbauer spectroscopy was also used to study the interactions between decamers and chitosan polymers with different Fe ions, concluding that there is an important participation of two NH₂ group and two OH from two glucosamine units. This result is in agreement with that one obtained by an X-ray study where a relation between two units of glucosamine by one metal ion salt molecule is proposed. The interaction of uranyl ions with chitosan was studied by UV-vis and FTIR spectroscopy; and the participation of only one NH₂ in the complex was proposed⁹. Domard ¹⁰ studied the copper chitosan complex by means of the circular dicroism and, in this case, the presence of the OH group in the complex besides the nitrogen atoms is postulated. In another work related to the interaction of Cu(II) with chitosan oligomers and the polymer dissolved using UV-vis, potenciometry and circular dicroism, the possibility to form complexes with the involvement of either one or two NH₂ is described.

EXPERIMENTAL PART

In the complex nomenclature, sometimes the amount of metal present is related to the polymer mass. This concentration is calculated by a mass balance between the amount of metal in the initial solution and the metal remaining in the final solution after the contact with the polymer. This determination is carried out by atomic absorption spectroscopy. For the specific case of mercury, the detection limit is 10 mg/L and the lower concentration is yet unknown. In this case, the “a” prefix in which the starting solution was 50 mg/L and the “b” prefix when the concentration was 600 mg/L was used.

In ESR spectroscopy, the complexes studied were prepared from metallic solutions with a 2000 mg/L concentration. With this technique, the influence of parameters such as degree of deacetylation and crystallinity of the polymer in the Cu(II) complex was studied. The influence of two different ion counters (NO₃ and Cl⁻) in the complex formation has also been studied.

Chitins crystalline index (C.I.) was determined by X-ray diffractogram, by means of the method described by Struszczzyk¹¹ for cellulose and applied to chitin through the following equation.

\[
CI = \left( \frac{I_{110} - I_{am}}{I_{110}} \right) \cdot 100
\]

Where \(I_{110}\) is the maximum intensity in the diffraction 110 and \(I_{am}\) is the intensity of the amorphous diffraction in the same unit \(\theta =12°30'\).

Chitosan crystalline index (C.I.) was determined by infrared spectroscopy from the absorbance relation \(A_{110}/A_{200}\) ¹². Polymers used in this case are:

High molecular weight chitosan of (Mₐ=300.000 g/mol) with C.I.=55.8% and deacetylation degree of 98% (QAPM-1). High molecular weight chitosan with (Mₐ=300.000 g/mol) previously dissolved and precipitated twice in order to decrease the crystalline (C.I.=46.26%) and deacetylation the degree remains at 98% (QAPM-2).

e-mail: galocardenas@udec.cl

Low molecular weight Chitosan (Mv=85,000 g/mol) with C.I.= 27.8% and deacetylation degree of 84% (QPBM-1).

α-chitin was obtained from squids, with a deacetylation degree of 16% and C.I.=82.7 

β-chitin was obtained from shrimps, with a deacetylation degree of 5% and C.I.=66.3 

Copper complexes were prepared by batch method using 100 mg of chitosan or its derivatives and 50 ml of copper nitrate p.a. from a 2000 mg/L solution.

ESR Methodology.

In this work, X band measurements ranges in a frequency zone from 9 to 10 GHz, and in a Q band, in which the microwave is around 34 GHz. A Bruker ESP380 model was used. As we expected, the band conditioned the magnetic field where the band appears. In the X-band, the signal corresponding to the free electron appears in the field between 320 mT and 350 mT, whiles the Q band and appears over 1.2 T. Besides, the spectrum analysis is analogous for the different bands.

One very peculiar characteristic of the ESR is that the spectrum is visualized in the first derivative, but the signals are detected as sensitive phases.

Table I: Values for the Hamiltonian spin parameters in the complex formed by the α and β-chitin with Cu(NO$_3$)$_2$ and CuCl$_2$.

<table>
<thead>
<tr>
<th>Copper Complex</th>
<th>Frequency (MHz)</th>
<th>$g_\parallel \pm \Delta g_\parallel$</th>
<th>$\Delta A_\parallel \pm \Delta A_\parallel$ (MHz)</th>
<th>$g_\perp \pm \Delta g_\perp$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chitin β-Cu(NO$_3$)$_2$ -X band</td>
<td>9831,396</td>
<td>2,33 ± 0.01</td>
<td>368 ± 46</td>
<td>2,07 ± 0.01</td>
</tr>
<tr>
<td>Chitin α-Cu(NO$_3$)$_2$ -X band</td>
<td>9831,161</td>
<td>2,32 ± 0.01</td>
<td>454 ± 46</td>
<td>2,07 ± 0.01</td>
</tr>
<tr>
<td>Chitin β-CuCl$_2$ -X band</td>
<td>9831,598</td>
<td>2,32 ± 0.01</td>
<td>418 ± 46</td>
<td>2,07 ± 0.01</td>
</tr>
<tr>
<td>Chitin α-CuCl$_2$ -X band</td>
<td>9831,237</td>
<td>2,32 ± 0.01</td>
<td>439 ± 46</td>
<td>2,07 ± 0.01</td>
</tr>
<tr>
<td>Chitin β-Cu(NO$_3$)$_2$ -Q band</td>
<td>34000</td>
<td>2,33 ± 0.01</td>
<td>480 ± 46</td>
<td>2,06 ± 0.01</td>
</tr>
<tr>
<td>Chitin α-Cu(NO$_3$)$_2$ -Q band</td>
<td>34000</td>
<td>2,32 ± 0.01</td>
<td>450 ± 46</td>
<td>2,07 ± 0.01</td>
</tr>
<tr>
<td>Chitin β-CuCl$_2$ -Q band</td>
<td>34000</td>
<td>2,32 ± 0.01</td>
<td>558 ± 46</td>
<td>2,07 ± 0.01</td>
</tr>
<tr>
<td>Chitin α-CuCl$_2$ -Q band</td>
<td>34000</td>
<td>2,32 ± 0.01</td>
<td>658 ± 46</td>
<td>2,07 ± 0.01</td>
</tr>
</tbody>
</table>

The signal corresponding to copper which values are in table I, exhibit a great apparent intensity in the α-chitin rather than in the β-chitin. This can be due to the fact that the α-conformation retains an amount of copper greater than the β-conformation. Besides, the α-complex is well defined and remains stable even if the NO$_3$ ion counter changes by the Cl presence. In the β-chitin, it does not occur in the same way, since the signal changes the form when the ion counter changes, showing a better defined signal corresponding to copper, when Cl was used. This observation suggests the participation of the ion counter in the complex formation.

The above analyses suggest that the complexes formed in each case are different, which is in agreement with the differences existed in the main arrangements. It is known that α-chitin can have a greater amount of OH groups closed one to another, because of the anti parallel chain arrangement. The interaction of these groups produces more rigidity than the β-chitin, where the arrangement is parallel and besides the OH is not enabled to interact.

This vicinity of the OH groups and the oxygen of the acetamide group in the α-chitin, proposes a possible coordination center for the metallic ions. It is expected that Cu(II) complexes are formed by the oxygen ligand of several chains. These complexes exhibit a low symmetry, because the metallic ions must get coordinated to the available sites, which are not abundant and showed great rigidity. The high values of $g_\parallel$ are according with these observations.

In a previous work, the presence of two types of metal coordination sites it was proposed and, in one of them, the nitrogen participate in the complex formation. Due to the great steric hinderance of the acetamide group, this is not able to get coordinated with the metal ion and the active site is related with the nitrogen, of the free amine group of the chitin. This is another way to find the deacetylation degree of the chitin related with the coordination site in the ESR spectrum. This type of site can appears in the chitosan and it has been mentioned above.

The α-chitin used, exhibit a deacetylation degree of 16 % and the β-chitin only 5%. As consequence, the α-chitin presents more free amine groups. However, for both chitin studied, the ESR presented only one coordination site. That site is different to that already mentioned and appears in the chitosan spectra. Then, it can be assumed that our observed coordination site can only be related to the oxygen atoms. The amine involved can be inhibited by the impurities presence.

RESULTS AND DISCUSSION

ESR Studies of Cu(II) complexes. Chitin – Cu(II) complexes

The polymorphic differences presented by chitin in the two more common forms (α and β), offers a different molecular environment for the metallic ions; each of them due to the different chain environment. Besides, the possibilities of coordination between metal ions such as Cu(II) and nitrogen can change in each case.

The ESR spectra obtained by means of MPC are shown in figures 1 and 2. In all of them, a typical signal of the Cu(II) axial complexes with a group of four peaks equally spaced, corresponding to the hyperfine parallel structure, and a perpendicular one can be observed. From this signal, Cu(II) center can be obtained besides the main parallel value of the hyperfine tensor. These parameters are summarized in table I.

In a few samples, another wide signal appears to be related to metallic impurities. In order to corroborate this statement, an ESR of the chitins without metal was measured and, as a matter of fact, they exhibit the presence of the same band found in the chitin-metal spectrum. The signal is quite wide and probably due to the presence of iron ions, mostly from the chitin. The high intensity of this band suggests that in the chitin, there is a greater amount of this impurity, rather than in the Cu(II) ion.
the chitosan QAPM-1 and QAPM-2, with the same deacetylation degree, the
same molecular weight and obtained from the same source, but with different
crystalline degree.

In the ESR spectrum presented in figures 3 and 4, it is observed that chit-
osan with lower crystalline degree (QAPM-2) produces wider bands and
with greater intensity than the chitosan more crystalline (QAPM-1), for both
ion counters. These characteristics of the bands can be correlated to a greater
amount of metal retention considering that all the other conditions remain con-
stant. From this analysis it can be concluded that metal ions are trapped in the
amorphous zone of the polymer.

It has been previously proposed that deacetylation degree possesses a great
influence on the amount of metal trapped by the polymer and it can be assumed
when the deacetylation degree increase, the amount of metal trapped increases.
This statement is corroborated from the ESR spectrum obtained in this work,
since the more deacetylated (QAPM-1 and QAPM-2, DD=98%), presents wider
bands and with greater intensity than chitosan lower deacetylated (QBPM-1,
DD=84%). QAPM-1, presents higher crystalline index than QBPM-1 and traps
a higher amount of metal, due to their higher DD.

**Table II:** Values of the Hamiltonian spin parameters in the chitosan com-
plexes with Cu(NO$_3$)$_2$.

<table>
<thead>
<tr>
<th>Copper complexes</th>
<th>Frequency (MHz)</th>
<th>$g_x \pm \Delta g_x$</th>
<th>$A_x \pm \Delta A_x$ (MHz)</th>
<th>$g_z \pm \Delta g_z$</th>
</tr>
</thead>
<tbody>
<tr>
<td>QAPM-1, Cu(NO$_3$)$_2$, X band</td>
<td>9831,076</td>
<td>2.264 ± 0.005</td>
<td>503 ± 22</td>
<td>2.091 ± 0.006</td>
</tr>
<tr>
<td>QAPM-2, Cu(NO$_3$)$_2$, X band</td>
<td>9831,467</td>
<td>2.261 ± 0.005</td>
<td>510 ± 22</td>
<td>2.089 ± 0.006</td>
</tr>
<tr>
<td>QBPM-1, Cu(NO$_3$)$_2$, X band</td>
<td>9831,539</td>
<td>2.261 ± 0.005</td>
<td>509 ± 22</td>
<td>2.091 ± 0.006</td>
</tr>
<tr>
<td>QAPM-1, Cu(NO$_3$)$_2$, Q band</td>
<td>34000</td>
<td>2.26 ± 0.01</td>
<td>491 ± 40</td>
<td>2.07 ± 0.01</td>
</tr>
<tr>
<td>QAPM-2, Cu(NO$_3$)$_2$, Q band</td>
<td>34000</td>
<td>2.26 ± 0.01</td>
<td>491 ± 40</td>
<td>2.07 ± 0.01</td>
</tr>
<tr>
<td>QBPM-1, Cu(NO$_3$)$_2$, Q band</td>
<td>34000</td>
<td>2.26 ± 0.01</td>
<td>491 ± 40</td>
<td>2.07 ± 0.01</td>
</tr>
</tbody>
</table>

It is very important to denote this phenomena when the salt of NO$_3$ salt
is changed by Cl$^-$. With the NO$_3^-$ one site type of retention is observed, on
the other hand, in the Cl$^-$ two types of sites can be observed. One of these sites
corresponds to the one observed in the NO$_3^-$ spectrum but the other is completely
different and besides, with the preference for the coordination of the metal ion
since their signal is more intense.

**Table III:** Values of the Hamiltonian spin parameters in the chitosan com-
plexes with CuCl$_2$.

<table>
<thead>
<tr>
<th>Copper complexes</th>
<th>Frequency (MHz)</th>
<th>$g_x \pm \Delta g_x$</th>
<th>$g_y \pm \Delta g_y$</th>
<th>$g_z \pm \Delta g_z$</th>
</tr>
</thead>
<tbody>
<tr>
<td>QAPM-1, CuCl$_2$, X band</td>
<td>9830,95</td>
<td>2.18 ± 0.01</td>
<td>--</td>
<td>2.07 ± 0.01</td>
</tr>
<tr>
<td>QAPM-2, CuCl$_2$, X band</td>
<td>9830,832</td>
<td>2.19 ± 0.01</td>
<td>--</td>
<td>2.04 ± 0.01</td>
</tr>
<tr>
<td>QBPM-1, CuCl$_2$, X band</td>
<td>9830,523</td>
<td>2.18 ± 0.01</td>
<td>--</td>
<td>2.08 ± 0.01</td>
</tr>
<tr>
<td>QAPM-1, CuCl$_2$, Q band</td>
<td>34000</td>
<td>2.17 ± 0.01</td>
<td>2.11 ± 0.01</td>
<td>2.05 ± 0.01</td>
</tr>
<tr>
<td>QAPM-2, CuCl$_2$, Q band</td>
<td>34000</td>
<td>2.18 ± 0.01</td>
<td>2.11 ± 0.01</td>
<td>2.05 ± 0.01</td>
</tr>
<tr>
<td>QBPM-1, CuCl$_2$, Q band</td>
<td>34000</td>
<td>2.17 ± 0.01</td>
<td>2.11 ± 0.01</td>
<td>2.05 ± 0.01</td>
</tr>
</tbody>
</table>

This new site, where the chlorine must be included inside the complex, col-
laps the hyperfine structure of copper and the axial system is converted into
a orthorhombic system where both main values of the tensor are very similar. This is observed more clearly in the Q band experiment (figure 5).

This phenomenon occurs when the system is formed by copper clusters where the ions are coupled by a long range of magnetic exchanges. In order to achieve this, the copper center must be formed by a short distance complex between them. The exchange produces a partial average of the tensor main value.

This clusters would be relatively long or very long chains (more than 8 Cu(II)) with common ligands, always in the same form and when the Cu(II) keeps the structure axis.

**CONCLUSIONS**

By using ESR, it is possible to characterize copper complexes with α-chitin and β-chitin, which are different in each case due to the difference in the existence of the chain arrangement. α-chitin retains an amount of copper greater than β-chitin. Besides, the complex with the α-chitin is well defined and remains, even when the NO₃⁻ ion counter has been changed to Cl⁻. In the β-chitin, the centers are different depending upon the ion counter.

The ESR allows the characterization of the copper complex with chitosan with different characteristics, such as crystallinity and degree of deacetylation and their influence in the complex formation is determined. The crystalline degree affects the capacity of chitosan to retain copper and, according to the experiments, the copper is trapped in a great amount when the polymer exhibits a lower crystalline index. The deacetylation degree also affects the amount of metal retention. A lower amount of acetyl groups increases the retention capacity. The linear copper clusters formation was also detected when Cl⁻ was used as ion counter. Finally, the symmetry of the metallic center depends on the original ion counter employed.

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