Silver Cation Coordination Study to AsW9 Ligand –
A Trilacunar Arsenotungstate Compound

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Objective: The main objective of this research is to find the coordination ratio between AsW9 and Ag+, as a preliminary study for synthesizing a new silver-arsenotungstate complex. Method and material: The ligand:cation molar ratio in complexes was determined by conductometric and potentiometric titrations of AsW9 with silver salts: CH3COOAg, AgNO3. Results: The ratio was obtained from the inflexion points of the curves when molar ratio was plotted versus conductivity, or from the equivalence point when silver added volume was plotted versus pH value. Each graphic shows one point of inflexion corresponding to 1:1.54 ratio of AsW9:Ag+. In the same manner, the equivalent volumes determined by graphical method gave the ratio 1:1.53. The spectral results confirmed that a AsW9:Ag+ complex was formed since the ligand absorption maxima values have been changed from 190 nm to 197 nm in the case of using AgNO3 and 196 nm for CH3COOAg corresponding to the W=O bond, and from 246.5 nm to 274 nm (AgNO3) and 270 nm (CH3COO Ag+) for the W-O-W bond. Conclusions: Silver cation exhibit a preference for AsW9 in a ratio of 3 to 2. This ratio can be associated to a sandwich type arrangement, with two trilacunar Keggin building blocks incorporating 3 metal cations in a tetrahedral geometry.

Keywords: arsenotungstate ligand, silver complexes, conductometry, potentiometric titration

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Introduction
Polyoxometalates (POMs) represent a large family of inorganic compounds with a wide range of applications, due to their redox activity, electronic density, structural and configurational diversity [1-4].

One of the widely-used polyoxometalate-ligand is the trilacunary Keggin unit. The Na3[HASW9O33]·17H2O (abbreviated as AsW9 inside the paper) used as ligand in this study, is an unsaturated arsenotungstate (which is a subclass of POMs family, with “As” as heteroatom and “W” as addenda) compound with a trilacunar Keggin structure. The electronic density of lacunae (one lacunae=six terminal oxygen) enables coordination of different metal cations [5-8].

Construction of multidimensional polyoxometalates structures using transitional metal cations as bounders have shown considerable interest in the research area over last years. New complexes combine intrinsic properties of each component as well as new properties arising from the combination of these components [9].

Among of the reported arsenotungstate complexes, the classical sandwich-type derivatives consist of two trilacunary Keggin building blocks, linked by 2 up to 6 transition metal cations at the equatorial plan [10-12].

Two different salts of the silver cation: AgNO3, which is widely used as a precursor for many silver complexes and CH3COOAg+ have been chosen. The silver ion is very versatile in creating supramolecular architectures due to its acceptor properties and the flexible coordination sphere. This metal ion can form from 2 up to 8 bonds with various donor atoms in a wide range of coordination geometries [13].

The most important oxidation state for silver in complexes is +1. The silver ion can adopt various coordination geometries such as linear (coordination number CN=2), the characteristic geometry explained by the ion’s tendency toward sd-hybridization), trigonal planar (CN=3), tetrahedral (CN=4, as in aqueous ion [Ag(H2O)4]+), octahedral (CN=6), even square antiprism (CN=8) geometry [13,14].

In POM complexes silver is generally found in tetrahedral coordinated geometry, being bounded by 2 terminal oxygen atoms from each lacunae of two different Keggin structures.

This paper is focusing on the study of AsW9 coordination with Ag+ as a preliminary data for synthesizing new complexes of this arsenotungstate ligand. The synthesis of POMs proceeds by self-assembly with a large variety of structures being obtained, depending on the reaction conditions, the nature and ratio of their constitutive elements, concentration, pH or T [7,9].

Methods
Materials and apparatus
AsW9 used in this study was synthesized as described earlier [15]. Metal cation salts (CH3COOAg and AgNO3) were of analytical grade from Alfa Aesar and Merck and were used without any further purification. All solutions were prepared by dissolving samples in purified water (obtained with NanoPure Diamond System (Barnstead, USA)) at a concentration of 10−3M for ligand (at a pH=7.5-8) and 2,5*10−3M for cations. UV spectra were recorded on an Analytik Jena SPECORD 210 spectrophotometer between λ = 190 - 400 nm in standard 1 cm quartz cuvettes. Con-
ductometric and potentiometric determinations were performed on an INOLAB 740 multiparameter device.

**Analysis method**

For every determination series, 40 mL of the ligand solution (10^{-5} M) was placed in the titration cell while the conductance and the pH of the solution was measured. Then, 4 μL of a 2,5\times10^{-2} M cation solution was added drop by drop in a stepwise manner using a calibrated micropipette. During titration conductivity and pH value were recorded after each addition. Titrations were made at 70-80°C. These values will be used in determination of ligand:cation coordination ratio. UV spectra were recorded before and after every titration to highlight modifications which occurred in each case.

**Results**

Results obtained by conductometry are presented in Figure 1. The graphic shows the conductivity values depending on the added titrant solution moles number.

In Figure 2 we have determined the equivalent volume for both salts by graphical method from potentiometric recorded data.

In Figure 3 we have plotted the overlaid recorded spectra of ligand and titrated solutions.

**Discussions**

By plotting molar ratio versus conductivity, the ligand:cation ratio was obtained from the inflexion points of the curves. In the same manner plotting pH versus added silver volume we have gathered the equivalence volume which exhibit a 2:3 ratio between ligand and cation. These ratios are presented in Table I. From the plotted data the same ligand:cation combination ratio can be noticed for both salts type.

The structure that correspond to this ratio is an arsenotungstate common sandwich-type structure. Each trilacunar Keggin unit has 6 terminal oxygen which can coordinate metal cations. There are 2 trilacunar Keggin units, one on the top of the other, with 3 silver cations between them, where each cation bound two terminal oxygen atoms per arsenotungstate ligand unit.

The spectral analysis confirm the formation of new complexes between ligand and cation due to a shifting of the peak corresponding to W=O_{bc}W bonds. The modified ligand absorption peak values are presented in Table II.

UV recorded spectra are characteristic to POM and show intense peak between 185 - 200 nm corresponding to W=O bonds. All spectra exhibit only the tail of this peak (Figure 3). This peak does not show large differences for W=O bonds in these compounds (all having the same addenda, W). The second peak is wider and is characteristic to W=O_{bc}W bonds (Table II). In complexes spectra, these second peaks move towards higher values of wavelength. This is due to the influence of coordinated cations especially on edges-tricentric bonds from AsW_9 structure [7, 16].

**Conclusions**

Silver cation exhibit a preference for AsW_9 in a ratio of 3 to 2 (from conductometry determinations) with a sandwich type arrangement. Two trilacunar Keggin building blocks incorporates 3 metal cations, each in a tetrahedral

![Fig. 1. Determination of inflexion point corresponding to molar ratio by conductometric titration](a) CH_3COO Ag, (b) AgNO_3

![Fig. 2. Determination of equivalent volume by potentiometric titration](a) CH_3COO Ag, (b) AgNO_3
geometry and each silver cation being bound by 2 terminal oxygen atoms from each Keggin structure. Data are confirmed by potentiometric analysis where the same ratio was noticed. These are preliminary data and further research will be performed to synthesize new arsenotungstate-silver complexes.

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Conflict of interest
None to declare.

References
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