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Preparation and characterization of colloidal copper xanthate nanoparticles

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Despite the important role of metal xanthates in a number of industrial processes and emerging applications, no attempts have been made to prepare the nanoparticles and to study colloidal solutions of insoluble heavy metal xanthates, in particular, copper. Here, we examined the formation of copper xanthate colloidal particles in reaction of aqueous solutions of cupric sulfate and various potassium xanthates, and characterized the products using UV-vis absorption, dynamic light scattering (DLS), zeta potential measurement, TEM, electron diffraction, FTIR, thermogravimetry, X-ray photoelectron spectroscopy (XPS), and X-ray absorption spectroscopy (XANES). Colloidal copper xanthates with ROCSSCu composition (R = ethyl, isopropyl, butyl, isobutyl, amyl groups), disordered structure and the average diameter of 40-80 nm easily formed and were stable for several hours, especially if excessive xanthate was used; the hydrodynamic diameter of nanoparticles was smaller at lower temperatures. TEM micrographs and S K-edge XANES revealed core/shell morphology of the particles, probably with Cu(I) bonded to four S atoms in the core and reduced copper coordination in the shell. Dixanthogens produced in the reaction along with ROCSSCu, seem to promote the nanoparticle aggregation and precipitate with the copper xanthate, affecting their thermal decomposition.

The structures and properties of bulk metal xanthates, including copper xanthates, and related substances, their soluble and adsorbed species have been reported in a number of studies (see, in particular, [3, 4, 6, 11, 20-31]). It is well known that the direct interaction of aqueous Cu2+ cations with xanthate ions yields a yellow precipitate of ROCS(S)Cu and dialkyl dixanthogen [20]

4ROCSSK + 2CuSO4 = (ROCSS)2 + 2ROCSSCu.

However, to the best of our knowledge, no studies have been aimed at the formation of the nanoparticles and colloids of xanthates of copper or other metal, which could be potentially used as carriers for poorly soluble metal xanthates in several applications, including flotation, biomedicine, materials science. Such a research is also important for understanding the behavior and role of aquatic metal xanthate nanoparticles formed, intentionally or unintentionally, in the flotation surreries, tail dumps and wastewaters of concentrating mills, water treatment systems. It is worth noting that the preparation of metal xanthate nanoparticles, especially for large-scale mineral processing and water treatment, should be easy and cost-effective.

In the current contribution, we describe the formation of colloidal copper xanthate particles via simple mixing of aqueous solutions of cupric sulfate and various potassium xanthates. It was demonstrated that the particles having ROCSSCu composition with an admixture of dixanthogen and devices [15-19].

Introduction

Metal xanthates, salts of the O-esters of carbonodithioic acids and the corresponding O,S-diesters ROCSSH, where stands for alkyl chain R (ethyl-, propyl-, butyl-, etc.), are widely used as collectors in the froth flotation of metal sulfide ores, reagents for heavy metal sedimentation in hydrometallurgy and wastewater treatment, in cellulose synthesis, as vulcanization accelerators in rubber industry, and some others [1-9]. Cytotoxic activity of metal xanthates on human cancer cell has been found [10, 11], however, testing of some compounds, including copper xanthates, has been retarded due to their low solubility [11]. In materials science and nanotechnology, xanthates have been proposed as capping agents for the synthesis of metal nanoparticles and self-assembled monolayers as alternative to thiols, and as sulfidizing reagents for preparation of metal sulfide nanoparticles [12-15]. The interest to metal xanthates as precursors of metal sulfide nanostructures rapidly grows in recent years since the low decomposition temperatures make them compatible with organic polymers for manufacturing of hybrid photovoltaic devices [15-19].
core/shell structure are largely amorphous, and their size and aggregative stability depend mainly upon the initial ratio of the reactants and temperature, the nature of alkyl groups is important too.

Experimental

Materials and preparation

Commercial reagents (about 95% purity) potassium ethyl xanthate (CH₃CH₂OCSSK), isopropyl xanthate (C₃H₇CHOCSK), n-butyl xanthate and isobutyl xanthate (C₄H₉CH₂OCSSK), and amyl xanthate (C₅H₁₁CH₂OCSSK) were recrystallized two times in acetonitrile and kept frozen in solid state; their fresh solutions were prepared using double distilled water directly before synthesis. Other chemicals were of analytical grade and were used as received. In a typical experiment of copper xanthate synthesis, 1 ml of CuSO₄ solution (0.3 mM) and 1 mL of potassium xanthate solution, which concentration varied from 0.15 mM to 12 mM in order to obtain a predetermined Cu to xanthate ratio, were added to 1 mL of water and agitated for 5 min at a desirable temperature (pH=8); the reaction mixture became yellow in several seconds. Then the solution was loaded into a cell for UV-vis absorption spectroscopy, dynamic light scattering (DLS), or zeta potential examination. The copper xanthate samples for spectroscopic examination (FTIR, XPS and X-ray absorption spectroscopy) and thermogravimetric analysis were precipitated from the 1 mM colloidal solutions or from more concentrated solutions with the same reagent ratios by centrifugation at 10 xg for 20 min, washed with water by decantation, and dried in air. Some samples were washed with acetonitrile in order to remove dianxathene.

Characterization

UV-vis absorption spectra were collected employing an Evolution 300 instrument (Thermo Scientific) in a thermostatic 1 cm quartz cell; when necessary, the solutions were diluted with water. DLS and zeta potential studies were conducted using Zetasizer Nano ZS spectrometer (Malvern Instruments Ltd, UK) at scattering angle 173° at a desired temperature in a folded polystyrene cell or polycarbonate cell with Pd electrodes. The volume weighted mean size (Zₚₜ) was normally considered in this study due to its high reliability.

Transmission electron microscopy experiments were performed with a JEM-2100 microscope (JEOL) operating at 200 kV. Samples were prepared by placing a hydrosol droplet on a carbon coated copper grid and allowing the solvent to evaporate at room temperature. FTIR spectra were recorded on a Bruker Vector 22 Fourier spectrometer in KBr pellets. Thermogravimetric (TG) and differential thermal analysis from ambient temperature to 900 °C was performed using STA449 F1 Jupiter instrument (Netzsch) at heating rate of 10 °C min⁻¹ in air or argon at a gas flow rate of 50 mL min⁻¹.

X-ray photoelectron spectra were collected from the precipitate attached to a conducting carbon tape with a SPECS instrument equipped with a PHOIBOS 150 MCD 9 hemispherical analyzer at electron take-off angle 90°. The monochromatized Al Kα excitation (1486.6 eV) was used for; the analytical chamber pressure was in the range of 10⁻⁵ mBar. The binding energies were corrected using the C 1s reference (285.0 eV) from aliphatic carbon; a low-energy electron flood gun was employed to eliminate inhomogeneous electrostatic charging of the samples. The spectra were fitted using Gaussian-Lorentzian peak profiles after Shirley background subtraction with CasaXPS software.

Cu K-edge and S K-edge X-ray absorption spectra were acquired at the KMC-1 dipole beamline (BESSY II synchrotron radiation facility, Helmholtz Zentrum Berlin) using the HIKE endstation with Si(111) crystal as monochromator simultaneously in total electron yield (TEY) and fluorescence yield (FY) modes. The measurements were carried out in a UHV chamber with a vacuum better than 1x10⁻⁷ mBar at room temperature.

Results and discussion

Fig. 1 shows typical electronic absorption spectra of initial xanthate solutions and the reaction media. The spectrum of xanthate displays absorption maxima A at 227 nm and C at 300 nm, which should be assigned to π-π* electron transition of C=S group and the π-π* transition of O-C (S)S moieties [22, 29, 30], respectively. These maxima decreased or disappeared, depending on the excess of xanthate in the reaction solution, and the new ones B at 256 nm originating from altered C=S bonding, and broad maxima D at 420-430 nm determining the
yellow color of the hydrosol from S – Cu bonds emerged. The transformations occurred in several seconds and then the spectra changed very little till precipitation of yellow copper xanthate took place after the time, usually few hours, depended on the mixture composition and temperature.

Representative TEM micrographs and selected area electron diffraction patterns (Fig. 2) demonstrate the formation of 30-80 nm particles tending to aggregation, at least upon drying the hydrosol, bridged possibly with residual dixanthogen. Closer examination of the images (see insertion) suggests that the nanoparticles are constructed of a denser core and about 5 nm thick shell. Broad diffuse rings were observed in electron diffraction pattern at approximately 0.12 nm and 0.2-0.25 nm; the first value is close to typical interatomic C-O and C-S distances in xanthate anion and the second one is typical for the alkyl chain length and Cu-S bonding [3, 25]. In general, the electron diffraction patterns are indicative of poor crystalline or amorphous structure of the particles. Precipitates from more concentrated solutions consist of mainly irregular 200-400 nm particles (right image).

Fig. 3 shows average hydrodynamic diameters ($Z_{av}$) and zeta-potentials of the reaction products for various xanthates after 15 min reaction as a function of their concentration. The size of the particles was about 120 nm in the media with the xanthate to copper ratios less than 0.5, passed maxima at the molar ROCSSK to CuSO$_4$ proportion of 0.5 and fell to less than 100 nm when excessive xanthates were added. The dimensions are in a reasonable accord with those observed using ex situ TEM. Zeta potentials were positive at the reactant ratios ≤0.5 and sharply decreased with increasing the content of xanthates. Such a behavior concurs with the above-mentioned reaction of Cu$^{2+}$ ions with two xanthate molecules [1-4, 20, 21], yielding copper xanthate and dixanthate. The positive surface charge of the particles prepared using stoichiometric quantities of cupric ions seems to induce the aggregation into micrometer entities while the excessive xanthate anions provide negative surface charge of the nanoparticles and their lesser hydrodynamic size. Generally, xanthates with smaller alkyl chains exhibit lower zeta-potential magnitude and form larger particles, especially in the copper-excessive media, and the situation is not straightforward at the xanthate to copper ratios above 0.5. Interestingly, the largest particles were obtained with ethyl xanthate. The mechanisms behind these phenomena are not quite clear; it cannot be ruled out that impurities remaining after xanthate purification or/and formed due to the reagent decomposition [1-4, 22] may play a role. It is worth noting that somewhat different and less reproducible results (not in Figures) were obtained in experiments with commercial xanthates, which were used as received; this should be taken into account when cheap commercial reagents are employed without additional purification, for example, in flotation practice.

The influence of reaction temperature and time on the particle size is illustrated in Fig. 4 for copper isobutyl xanthate. One can see that the differences in the hydrodynamic diameter arose in first seconds of the reaction, and the particles grow faster at higher temperatures. Usually, the effect of temperature was less pronounced if more extra xanthate was added, suggesting that aggregation of nanoparticles is at least one of the reasons of the particle growth.

Thermogravimetric analysis of the copper butyl xanthate as an example (Fig. 5) showed that the sample separated by centrifugation lost ~75 mass.% at about 190 °C, and the sample treated then with acetone for dixanthogen removing lost about 60 mass.% at the temperature by 10 degree higher. The main weight difference was due to evaporation of about 15 mass.% of the dixanthogen admixture at 160-170 °C, both in air and under inert atmosphere. The copper sulfides produced by the xanthate decomposition, which exhibited somewhat behavior in air between 260 °C and 800 °C (not shown in the Fig. 5), require further investigation that is beyond the framework of the current article.

FTIR spectra (Fig. 6) of the samples obtained from the solutions with various concentrations and ratios of reagents, including those with excessive Cu$^{2+}$ ions, were very similar, indicating that the particles have the same bulk composition and structure, and their diverse behavior in solutions (Fig. 3) is
due to interfacial phenomena. The spectra from the copper xanthates, and dixanthogen and potassium xanthate as reference materials coincide with those reported by Little and co-workers [21] and confirm the presence of dixanthogen in the precipitated samples.

Dixanthogens evaporated under ultra-high vacuum conditions of XPS experiment, and the elemental concentrations of the copper xanthates derived from the photoelectron spectra (in particular, for copper n-butyl xanthate, at. %: Cu 8.5, S 17, O 14.8, C 59.6) approaches the Cu to S ratio of 1 to 2 and so the ROCSSCu composition, taking in mind an inevitable surface contamination by carbon and oxygen-bearing species. The Cu 2p spectra (Fig. 7) with Cu 2p3/2 peak at 932.9 eV, no shake-up satellites at 944-948 eV, and Auger Cu LMM peak position at 916 eV mean that copper occurs as Cu(I) [32-36]. The S 2p band can be fitted using the principal component with the S 2p3/2 binding energy at 162.3 eV and a minor one at 164.3 eV (about 5 relat. %); the latter may be attributed to bridging sulfur atoms in residual dixanthogen [36, 37] uptaken by the disordered cuprous xanthate. The C 1s spectrum diverges from that of potassium xanthate, which contains the lines of aliphatic carbon at 285.0 eV, and equal intensity of the bands at about 286.5 eV (R-C-) and 288 eV (-O-C-S-) [36, 37], possibly, because the xanthate radical in the surface layers of the particles is oriented outward or due to adsorption of products of xanthate oxidation (particularly, butanol) and carbon contaminants, and some copper xanthate surface decay in vacuum under X-ray irradiation. Akin to FTIR, the photoelectron spectra acquired from the samples prepared with an excess or deficit of butyl xanthate and at various concentrations of the reagents were essentially the same, and, on the whole, agreed with those reported for cuprous xanthate anions. Dixanthogen, which is another main reaction product poorly soluble in water, forms droplets charged negatively (to be published in detail elsewhere), which should interact more strongly with copper xanthate particles positively charged in the media with excessive copper ions than with negatively charged ones in the presence of excessive xanthate anions. Dixanthogen partially co-precipitates together with copper xanthate, covering and bridging the nanoparticles and affecting their properties. Up to a few per cent of dixanthogen may be incorporated in the copper xanthate particles, disordering their structure. It is interesting, in particular, that the copper butyl xanthate precipitate containing some dixanthogen decomposes at a lower temperature than that after extraction of dixanthogen.

The phenomena and species described above should play a role in the transport of copper xanthates in the environment, technological or biological media. The findings should be taken into account or implemented in several fields, including flotation practice, water treatment, manufacturing of copper xanthate and copper sulfide nanostructures for various applications.
Conclusions

It has been demonstrated for the first time that colloidal copper (I) xanthate nanoparticles having the composition ROCSSCu (R = ethyl, isopropyl, butyl, amyl groups) and the diameter of 40-80 nm can be simply formed by mixing of aqueous solutions of cupric sulfate and potassium xanthates. The UV-vis spectra evidenced fast appearance of absorption maximum at about 430 nm, suggested that the reaction was largely completed in less than 1 min. A sharp change in zeta potential and hydrodynamic diameter of the particles occurred at the xanthate to cupric ion molar ratio of 0.5 corresponding to the reaction stoichiometry. The particles produced using excessive xanthate concentration had high negative zeta potential magnitude, lower diameter and higher stability than the ones charged positively at the excess of cupric ions. The smaller particles emerged and grew more slowly at lower temperatures. Characterization of copper xanthates prepared using various concentrations and ratios of the reagents found that the particles had very similar composition. IR spectroscopy and thermogravimetric analysis demonstrated that the precipitated material comprised dixanthogen (an order of 15 mass. %), which possibly promoted aggregation of the colloidal copper xanthate, especially in the solutions with excessive copper ions, and decreased the temperature of thermal decomposition of the precipitates. XPS data suggested that minor dixanthogen may be incorporated in the disordered particles. TEM micrographs, electron diffraction and S K- and Cu K-edge XANES spectra collected in TEY and FY modes revealed a core/shell structure of the particles, showing probably reduced coordination and somewhat increased local positive charge of Cu atoms in the shell in comparison with the copper in a distorted tetrahedral coordination with four S atoms in the core.

Acknowledgements

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Colloids formed via direct interaction of aqueous Cu\(^{2+}\) ions and xanthates and their precipitates incorporating dixanthogen were characterized.