Spectral Study of Silver Nanoparticles Prepared by Chemical Method

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ABSTRACT

Silver nanoparticles were prepared by the reduction of silver salt (silver nitrate AgNO₃) solution using three different methods. The main differences between these methods is by belong to the reducing agent. It is found that the prepared silver nanoparticle was pure and stable for long time. UV-VIS spectrometry indicated formation of spherical silver nano particles. The surface plasmon resonance peaks in absorption spectra for silver colloidal solution showed that the absorption maximum range was at 380- 420 nm. The structures are confirmed by X-ray diffraction (XRD) and the crystallite size was determined from X-ray line broadening using the Scherrer's equation and it was about 32 nm.AFM measurements show that silver nano particles have the average diameter of 69.39 nm and 81.91 nm.

الدراسة الطيفيه لدقائق الفضةالنانويةالمحضره بالطريقة الكيميائية

الخلاصه

تم تحضير دقائقالفضه النانوية باستخدام ثلاثة طرق مختلفه وذلك باختزال نترات الفضة (AgNO₃).ان الاختلاف الرئيسي بين هذه الطرق هو استخدام العامل المختزل. دقائق الفضه النانويه المحضرة وجدت نقيه ومستقرّة لوقت الطويل.

جهاز قياس الطيف اشار إلى تشكيل الدقائق النانوية الفضية الكرويه. قمم طيف الامتصاص للمحلول الفضّي اظهرت بأنّ الإمتصاص الأعظم كان في مدى(1800- 380).

اظهرت بأن الإمتصاص الأعظم كان في مدى(-nm420). التركيب لدقائق الفضه النانويه والحجم البلوري مؤكّدة بوسطة قياس حيود الاشعة السينية (XRD) و بإستعمال معادلة Scherrer's كان حوالي 32nm.

INTRODUCTION

olloidal particles are increasingly receiving attention as important starting points for the generation of micro and nano structures [1]. These particles are under active research because they posses interesting physical properties differing considerably from that of the bulk phase. It comes from small sizes and high surface/volume ratio [2]. Silver nano particles have received considerable attention due to their attractive physical and chemical properties. Metallic silver colloids were first prepared more than a century ago. Ag nanoparticles can be synthesized using various methods: chemical, electrochemical [3], γ -radiation [4] photochemical [5], laser ablation

[6] etc. The most popular preparation of Ag colloids is chemical reduction of silver salts by sodium borohydride, sodium citrate and cetyltrimethlyammonium bromide, (CTAB). This preparation is simple, but the great care must be exercised to make stable and reproducible colloid. The purity of water and reagents, cleanliness of the glassware are critical parameters. Solution temperature, concentrations of the metal salt and reducing agent, reaction time influences particle size. Controlling size and shape of metal nanoparticles remains a challenge [7]. The size-induced properties of nanoparticles enable the development of new applications or the addition of flexibility toexisting systems in many areas, such as catalysis, optics, microelectronics and so on. Silver nanoparticles exhibit new optical properties, which are observed neither in molecules nor in bulkmetals. One example is presence of absorption band invisible light region. This band appears due to the surface Plasmon oscillation modes of conduction electrons which are coupled through the surface to external electromagnetic fields [8]. The surface plasmon resonance and large effective scattering cross section of individual silver nanoparticles make them ideal candidates for molecular labeling, where phenomena such as surface enhance Raman scattering (SERS) can be exploited [9]. In addition, silver nanoparticles have recently been shown to be apromising microbial material [10]. In this research, Ag nanoparticles were synthesized by silver salt (AgNO3) reduction with sodium citrate, sodium borohydride and cetyltrimethlyammonium bromide (CTAB).

EXPERIMENTAL

Preparation of Ag colloid

Silver nitrate with trisodium citrate

The materials used in this method silver nitrate (AgNO₃) and trisodium citrate (TSC) ($C_6H_5O_7Na_3$) of analytical grade purity. The silver colloid was prepared by using chemical reduction method according to the description of Lee and Meisel [11].

All solutions of reacting materials were prepared in distilled and deionized water (DW). In a typical experiment 25 ml of 0.01g (AgNO₃) was heated to boiling. To this solution 20 ml of 0.025 gtrisodium citrate was added drop wise. The solution was mixed vigorously, heated until color's change is evident (pale yellow). Then the solution was cooled to room temperature while it is stirring. The Ag nanoparticles prepared by this method have low stability for short time.

The chemical for such reduction can be written as follows:

$$4Ag^{+} + C_{6}H_{5}O_{7}Na_{3} + 2H_{2}O \rightarrow 4Ag^{0} + C_{6}H_{5}O_{7}H_{3} + 3Na^{+} + H^{+} + O_{2}\uparrow$$

Silver nitrate with oxalic acid

Colloidal Ag nanoparticles were prepared by reducing silver nitrate with oxalic acid in presence of (CTAB). Three solutions were prepared. First dissolve 0.01 g of AgNO₃ in 5ml DW second, dissolve 0.01 g of CTAB in 10 ml of DW third, and dissolve 0.05 g of oxalic acid in 10mlDW. The solution of CTAB was addeddropwise to the solution of nitrate. After that dropwise the oxalic acid solution to the results of CTAB and nitrate. After mixing oxalic solution, the color result to change light acorned followed that, the color change to dark acorned. Finelly, the color of the result solution become yellow.

Silver nitrate with sodiumborohydride

This method based on mixing silver nitrate with (NaBH₄). Firstly dissolving 0.0001 g of AgNO₃ in 5 ml ofDW and0.0006 g of NaBH₄ in 5ml DW. The solution ofNaBH₄become light yellow after dropwise of AgNO₃ solution with storing stirring. If the stirring is continued once all the silver nitrate has been added, aggregation is likely to occur, the yellow darkens, turns violet.

The chemical equation for this method can be written as follows:

 $AgNO_3 + NaBH_4 \rightarrow Ag+1/2H_2+1/2B_2H_6 + NaNO_3$ characterizations of NPs

-Absorbance spectra (SPE spectra) of NPs solution were measured by UV-VIS double beam spectrophotometers, CECIL C. 7200 (France) and SHIMADZU. All spectra were measured at room-temperature in a quartz cell with 1 cm optical path.

The Agnanoparticles samples were studied by X-ray diffraction technique (Philips PAN analytical, The Netherland) using Cu K α radiation. The measure condition X-ray target Cu, wavelength 15.406 (nm), generator voltage and current was set at 40 KV and 30 mA respectively. The Ag nanoparticles samples were scanned in the 20 range 20 to 70 (deg) ranges in continuous scan mode. The scan rate was 5 deg/sec.

The AFM analysis was done using scanning prop microscopy (CSPM-3000). It is an instrument capable of measuring the topography of a givensample. A nanosized tip attached on a cantilever is traded over the sampleand a 3D image of the sample topography is generated on a computer.

RESULTS AND DISCUSSIONS

UV-VIS spectroscopy analyses

Figure (1) Shows UV-VIS spectroscopy of Ag nanoparticles produced using mixing AgNO₃and TSC. Which indicates that the surface plasmon resonance peak was broad and located between 350 to 720 nm with an absorption maximum at 450 nm. Comparison of the experimental results with reference results [12] which showed the diameter range of AgNPs in colloidal solution is between 75 and 110 nm.

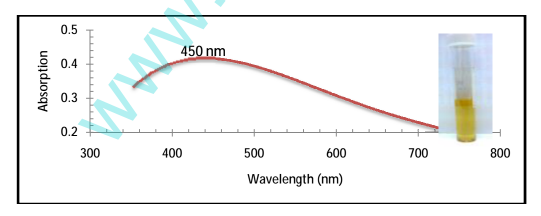


Figure (1)UV-VIS spectral and photo of Ag nanoparticles by chemical reduction AgNO₃ with TSC.

Figure (2) shows the color of colloidal Ag nanoparticles product using of AgNO₃with CTAB and oxalic acid. Figure (2). Depicts 4 color belong to colloidal Ag nanoparticles with time of storage. It can be concluded that, the reducing AgNO₃ used in this method was slow, therefore, the reaction needs to more time to complete. The reducing agent reduces large Ag nanoparticles to form small Ag nanoparticles after period of time.



Figure (2) photo shows formation stages Ag nanoparticles prepared by AgNO₃ with CTAB with oxalic acid.

Figure (3) represents the UV-VIS spectroscopy of colloidal Ag nanoparticles produced AgNO₃, CTAB, and oxalic acid. The peak of surface Plasmon resonance appears at 407 nm with peak width half maximum (PWHM) at 95 nm. After that the peak of SPR be broaden. The increase of peak height indicates increases of Ag nanoparticles concentration. The Ag nanoparticles prepared in this method have high stability with long time.

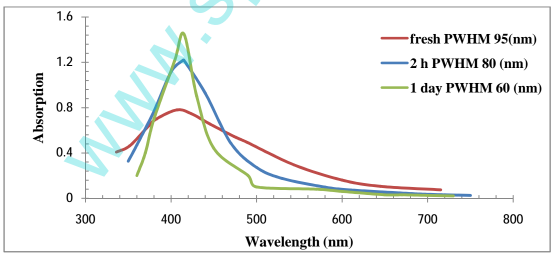


Figure (3) UV-visible spectra of formation stages Ag nanoparticles prepared by AgNO₃ with CTAB with oxalic acid.

Figure (4) Shows that the UV-VIS spectroscopy colloidalAgNPsusingAgNO₃, NaBH₄. The preparation of Ag nanoparticles have low stable for short time, because the drawbacks to this synthesis were that the sodium borohydride was unstable. The nanoparticles will be aggregated, although this can be avoided by capping the nanoparticles with some stabilizing agent such as polyvinyl alcohol .The surface Plasmon resonance peak absorption maximum at 386.6 nm as show in Figure (4).

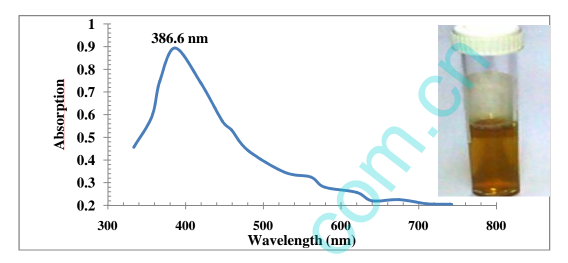


Figure (4) UV-VIS spectral and photo of yellow Ag nanoparticles was produced by chemical reduction of AgNO₃with NaBH₄.

X-RAY Diffraction analyses

Figure (5)Shows a typical X-ray diffraction (XRD) of the collidal Ag nanoparticles prepared by using NaBH₄ as a reducer. The reflection peak can be indexed to face-centered cubic silver. The lattice constant (a) calculated from this pattern is 4.085°A, which is consistent with the standard value of 4.085 °A (JCPDS Card File 04-0783). No impurities are detected from this pattern. This indicates that pure silver metal is obtained under the present synthetic conditions. The crystallite size of the silver nanoparticles equal 32 nm. It was determined from X-ray line broadening using the Scherrer's equation.

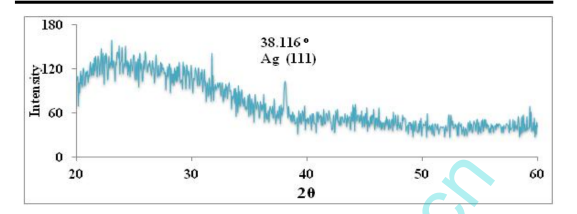


Figure (5)X-Ray diffraction of Ag nanoparticles was produced by chemical reduction AgNO₃with NaBH₄.

Atomic Force Microscope (AFM) analyses

Figure (6) Shows AFM images and corresponding size distributions of Ag nanoparticles were prepared by reducing AgNO₃ by (CTAB) in presence of oxalic acid. The Ag nanoparticles were storage at room temperature for eight months before carrying out AFM measurement, without addition of any stabilizer agent. The nanoparticles thus prepared were calculated to have the average diameter of 69.39 nm.

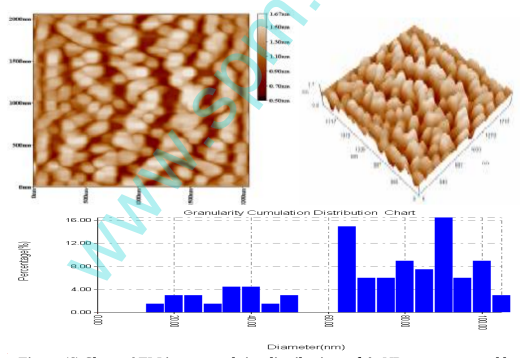


Figure (6) Shows AFM images and size distributions of AgNPswere prepared by reducing silver nitrate with (CTAB) in presence oxalic acid.

Figure (7) Shows AFM images and corresponding size distributions of Ag nanoparticleswere prepared by reducing $AgNO_3$ with $NaBH_4$. The silver nanoparticles were storage at room temperature for eight months before carrying out AFM measurement, without addition of any stabilizer agent. The nanoparticles thus prepared were calculated to have the average diameter of 81.91nm.

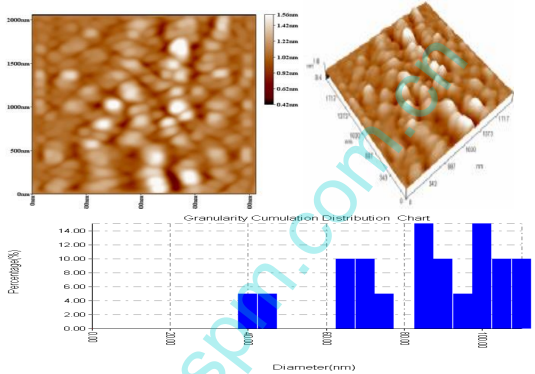


Figure (7) Shows AFM images and size distributions of AgNPswere Prepared by reducing AgNO₃ with NaBH₄.

CONCLUSIONS

The colloidal Ag nanoparticles prepared using three different method depends on the change of reducing agent (TSC, CTAB and NaBH₄) with AgNO₃. On the other hand, the effect of oxalic acid are studied on the formation of Ag nanoparticles by UV-VIS spectroscopy result.UV-VIS spectroscopy indicated formation of spherical silver nanoparticles. The surface plasmon resonance peaks in absorption spectra for silver colloidal solution showed that the absorption maximum range was at 380- 420 nm. X-ray measurementshave shown that the crystallite size of the AgNPs equal 32 nm. It was determined from X-ray line broadening using the Scherrer's equation.AFM measurements show that AgNPs have the average diameter of 69.39 nm and 81.91 nmand tend to form aggregates.

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