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ORIGINAL RESEARCH

One-step, low-temperature fabrication of CdS quantum dots by watermelon rind: a green approach

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http://dx.doi.org/10.2147/IJN.S79988

Abstract: We investigated the one-step synthesis of CdS nanoparticles via green synthesis that used aqueous extract of watermelon rind as a capping and stabilizing agent. Preliminary phytochemical analysis depicted the presence of carbohydrates which can act as capping and stabilizing agents. Synthesized CdS nanoparticles were characterized using UV-visible, Fourier transform infrared spectroscopy, X-ray diffraction, EDX, dynamic light scattering, transmission electron microscopy, and atomic force microscopy techniques. The CdS nanoparticles were found to be size- and shape-controlled and were stable even after 3 months of synthesis. The results suggest that watermelon rind, an agro-waste, can be used for synthesis of CdS nanoparticles without any addition of stabilizing and capping agents.

Keywords: green synthesis, nanoparticles, biomimetic, atomic force microscopy

Introduction

Cadmium sulfide (CdS) nanoparticles (NPs) are typical II-VI semiconductors with unique optical properties and tunable photoluminescence compared to their bulk counterparts on account of the size quantization effects. The preparation of CdS NPs has been carried out using various methods such as microwave heating,¹ microemulsion synthesis,² chemical synthesis,³ and ultrasonic irradiation.⁴ To fulfill the growing need for environmentally friendly NPs, researchers are using microorganisms for synthesis of various metal NPs.⁵ Nowadays, plant and agriculture waste extracts are used as reducing and capping agents for the synthesis of NPs. An extensive literature survey revealed that there are no reports on the synthesis of CdS NPs using agricultural wastes. Agricultural wastes are known to be rich in organic molecules and biopolymers which can play versatile roles in various areas. Agro-wastes and industrial byproducts are explored for a variety of applications, such as adsorption⁶ and corrosion and as precursors for activated carbon. Recently, the ability or potential of agro-wastes toward synthesis of noble NPs is being explored. Interestingly, fruit residues and wastes have proved to be good stabilizers for fabrication of NPs. However, the ability of fruit residues and wastes for the fabrication of metal sulfides and oxides are yet to be explored systematically. In view of this, the present study reports the feasibility of fabrication of CdS NPs by watermelon rind (WR), an agro-waste produced at high rates during summer.

WR consists of pectin, citrulline, cellulose, proteins, and carotenoids.⁷ These polymers are rich in functional groups such as hydroxyl (cellulose), amine (proteins), and carboxylic (pectin) that can easily bind inorganic materials. Recently, WR was reported to bind Ni²⁺ and Co²⁺ ions from aqueous solution.⁸

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International Journal of Nanomedicine 2015:10 (Suppl I: Challenges in biomaterials research) 183–188 **183** © 0.05 (Suppl I: Challenges in biomaterials research) 183–188 **183** © 2015 Lakshnipathy et al. This work is published by Dove Medical Press Limited, and licensed under Creative Commons Attribution – Non Commercial (unported, v3.0) permission from Dove Medical Press Limited, provided the work is properly attributed. Permissions beyond the scope of the License are administered by Dove Medical Press Limited, Information on how to request permission may be found at http://www.dovepress.com/permission.mayb This innovative research reports a one-step, simple approach for the "green" synthesis of CdS NPs using WR, an agro-waste. The phase, morphological, and elemental analysis of the as-synthesized CdS NPs are investigated.

Materials and methods Preparation of extract

WRs were obtained from a local fruit market, washed with tap water, and dried under sunlight for 7 days. The dried WR pieces were washed repeatedly with hot water (70°C) to remove any soluble matter present and then dried in an oven at 85°C for 48 hours. To 50 mL of distilled water, 1 g of sieved WR was added, and the mixture was boiled in a water bath for 30 minutes to prepare an aqueous extract. The extract was then cooled to room temperature and filtered through 0.45 μ m filters, and this freshly prepared extract was used for synthesis. This aqueous extract was termed as WR aqueous extract (WRAE).

Synthesis of CdS NPs

To 20 mL of 0.25 M cadmium chloride (CdCl₂) solution, 5 mL of sodium sulfide (Na₂S) was added until it attained an orange-yellow color indicating the formation of Cds; 10 mL of freshly prepared WRAE were added to the colored solution and stirred for 12 hours at 70°C on a magnetic stirrer. The colloidal solution was centrifuged at 10,000 rpm for 10 minutes, and the obtained orange-yellow solid was washed with water, methanol, and acetone to remove biomolecules. The solid was then dried at 85°C and used for further characterization. The synthesized CdS NPs were characterized using UV-visible spectroscopy, X-ray diffraction (XRD), Fourier transform infrared spectroscopy (FTIR), dynamic light scattering (DLS), atomic force microscopy (AFM), and energy-dispersive X-ray spectroscopy (EDX) analyses.

Analytical techniques

The sample aliquots from different concentrations of extracts were monitored using a Hitachi double-beam UV-visible spectrometer (model: U-2800) in the range of 200-800 nm at a scan speed of 400 nm/min with a resolution of 1 nm. FTIR analysis was done using a Thermo Nicolet Avatar 330 FTIR for watermelon rind powder (WRP) and dried WRAE in the range of 4,000–400 cm⁻¹ to determine the functional groups bound onto the CdS NPs. The powder XRD analysis of ovendried CdS NPs was carried out with a Bruker D8 Advance model equipped with a Lynx eye detector and beta Ni filter. The scanning was done in the range of 2θ from 10° to 80° at 0.02°/min. The size and morphology of CdS NPs synthesized by WR extract were examined using a transmission electron microscope (Philips CM200 model). The samples were prepared on carbon-coated copper grids prior to the transmission electron microscopy (TEM) analysis. The particle size analysis was carried out by DLS (DelsaTM Nano).

Results

The WRAE was initially evaluated for identification of a capping agent responsible for synthesis of CdS NPs. The preliminary phytochemical analysis of the extract confirmed the presence of carbohydrates through Molisch's test by the formation of purple-colored ring with addition of concentrated H₂SO₄. Previous researchers have reported starch to be an effective capping agent for green synthesis of CdS and CdSe NPs.^{9,10} But in the present case, water-soluble fractions of WR such as carbohydrates might act as a capping agent for synthesis of CdS NPs. The hydroxyl group of carbohydrates act as passivation contacts for the stabilization of the CdS NPs in the aqueous solution.¹¹ A schematic representation of passivation of carbohydrates for stabilization of CdS NPs in aqueous solution is shown in Figure 1.

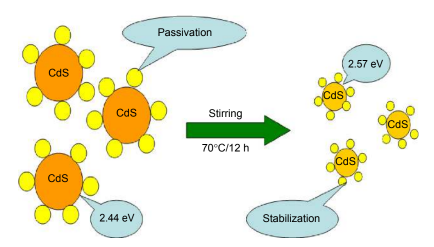


Figure I Plausible mechanism of CdS nanoparticles stabilized by watermelon rind aqueous extract.

FTIR spectroscopy was used to analyze the type of organic capping agent adsorbed onto the surface of synthesized CdS NPs. The FTIR spectrum of synthesized CdS NPs shows strong bands at 3,416, 3,139, 2,924, 1,619, 1,400, 1,122 and 1,067 cm⁻¹ (Figure 2A). The bands at 3,416 and 1,619 cm⁻¹ correspond to symmetric and asymmetric stretching vibrations of -OH groups. Bands at 3,139 and 2,924 cm⁻¹ correspond to aromatic and aliphatic -CH stretching vibrations. A peak at 1,400 cm⁻¹ corresponds to –CH bending vibrations and peaks at 1,122 and 1,067 cm⁻¹ are due to symmetrical and asymmetrical bending vibrations of aromatic -CH groups. It has been reported that CdS can give a very weak absorption band at 405 cm⁻¹.¹² A very weak band was observed at 405 cm⁻¹, which corresponds to CdS NPs (Figure 2B). The bands from the FTIR spectra confirmed that carbohydrates were present on the surface of the CdS NPs as capping and stabilizing agents.

The formation of CdS NPs was monitored by UV-visible spectroscopy. An absorption maximum at 484 nm with a band gap of 2.57 eV was observed, which was due to the quantum size effect of formed CdS NPs.¹³ The band gap of bulk CdS before addition of WRAE was found to be 2.44 eV; this could be due to the consequence of a size quantization effect in the sample.¹⁴ The extract concentration was varied in order to study the effect on size confinement. It was observed that the highest extract concentration resulted in size-controlled CdS NPs. The results were confirmed by DLS analysis. In order to determine the elemental composition of synthesized NPs, the EDX pattern was recorded. The EDX pattern of synthesized

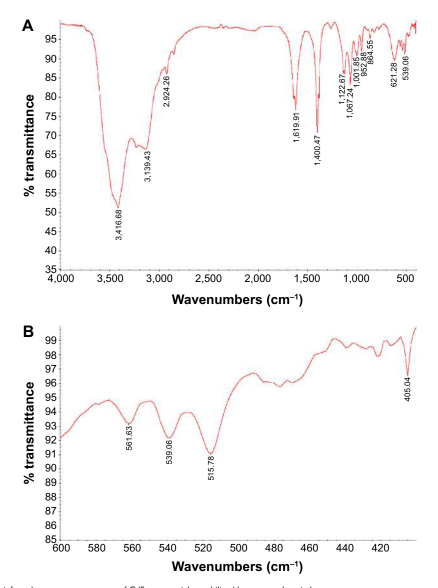


Figure 2 Fourier transform infrared spectroscopy spectra of CdS nanoparticles stabilized by watermelon rind aqueous extract. Notes: (A) Full range. (B) Depiction of CdS peak.

185

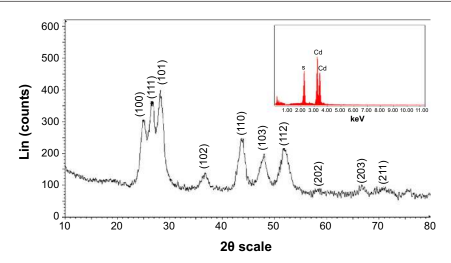


Figure 3 X-ray diffraction and energy-dispersive X-ray spectroscopy (inset) patterns of CdS nanoparticles fabricated using watermelon rind aqueous extract.

NPs showed strong signals of Cd and S alone (Figure 3 [inset]), without contamination by other atoms.

XRD analysis was done for the WRAE-synthesized CdS NPs to confirm the crystalline nature (Figure 3). The XRD patterns showed the presence of broad peaks at 20 values of 24.90, 26.53, 28.14, 43.63, 47.99, and 51.75, respectively, which is attributed to the (100), (111), (101), (110), (103), and (112) diffraction planes of hexagonal-structure CdS NPs, corresponding to the JCPDS number 80-0006. The broad peaks indicate the particles were of very small crystallite size. The mean crystallite size was calculated to be 28 nm using the Scherrer formula.¹⁵ The results indicated that WRE acted as a stabilizing and capping agent in the preparation of the hexagonal CdS NPs.

DLS analysis (Figure 4) was performed in order to determine the particle size distribution of the synthesized CdS NPs mediated by WRAE. It can be seen from Figure 4 that around 94% of CdS NPs were 90 nm in size. The particle sizes of CdS NPs determined using DLS were corroborated

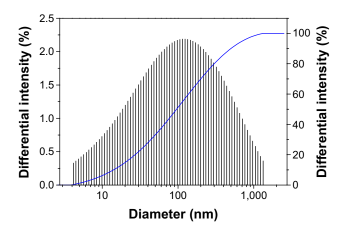


Figure 4 Particle size plot of synthesized CdS nanoparticles stabilized by watermelon rind aqueous extract.

with AFM analysis. The size and shape of CdS NPs were found to be stable even after 3 months of synthesis. TEM analysis was carried out in order to determine the shape of the as-synthesized CdS NPs.

A TEM image of CdS NPs is shown in Figure 5. The image indicates CdS NPs were shape controlled with a diameter range of 90 nm. This was confirmed by the XRD and AFM analyses.

Figure 6 shows a typical AFM image (Easyscan 2; Nanosurf, Liestal, Switzerland) of CdS NPs stabilized by WR. Most of the NPs were found to range in size between 80 and 90 nm, which is larger than the values calculated from the XRD pattern.

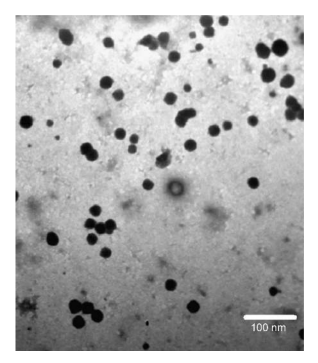


Figure 5 Transmission electron microscopy image of watermelon rind aqueous extract-stabilized CdS nanoparticles.

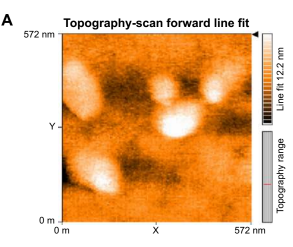


Figure 6 Atomic force microscopy image of synthesized CdS nanoparticles. Notes: (A) Topography-scan forward line fit and (B) deflection-scan forward line fit.

The larger particles from the AFM image can be ascribed to the aggregation of the smaller particles.¹⁶ The mean surface roughness was measured and calculated to be 16 nm by AFM, which indicates that relatively uniform CdS NPs were synthesized by the one-step green process.

Conclusion

One-pot green synthesis of CdS NPs was successfully performed using the aqueous extract of WR, an agro-waste. Water-soluble carbohydrates present in WRAE were believed to be the capping and stabilizing agent for the CdS NP formation. The capping of biomolecules onto CdS NPs from WRAE was confirmed by FTIR analysis. The average particle size of CdS NPs was found to be around 90 nm. The shapes of the particles were spherical, confirmed by TEM analysis. The surface roughness was measured and calculated to be 16 nm by AFM. Thus, it is proved that WR, an agro-waste, has been effectively used as a capping and stabilizing agent for synthesis of CdS NPs using a green chemistry approach.

Acknowledgments

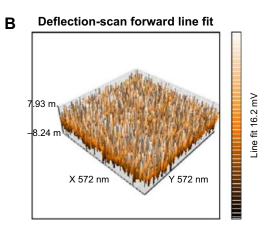
The authors gratefully acknowledge the management of VIT University, Vellore, India for encouragment, and providing the necessary facilities for carrying out this work, and we thank Johnson & Johnson for the publication sponsorship.

Disclosure

The authors report no conflicts of interest in this work.

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