

C. r. er. li . a ailable a SciVe e Scier.ceDi ec

Materials Letters

, real h. me age: .el. e ie .c. m/l. ca e/ma le



Facile s nthesis of stable cadmium sulfide quantum dots with good photocatal tic activities under stabili ation of h drophobic amino acids

Yuming Guo*, Lili Jiang, Lingling Wang, Xiaoman Shi, Qilong Fang, Lin Yang*, Fangfang Dong, Chuangchuang Shan

College of Chemistry and Environmental Science, Engineering Technology Research Center of Motive Power and Key Materials of Henan Province, Henan Normal University, Xinxiang 453007, PR China

ARTICLE INFO

Article history: Received 21 November 2011 Accepted 13 Januar 2012 Available online 21 Januar 2012

Keywords: Cadmium sulfide Quantum dots H drophobic amino acids Photocatal sis

ABSTRACT

In the present stud, the stable CdS quantum dots were successfull prepared using h drophobic amino acids methionine and valine as stabili ers through a facile one-pot strateg in aqueous media. The CdS quantum dots e hibited strong absorption and photoluminescence properties in the visible wavelength region. From the results, the CdS quantum dots can effectivel photocatal ticall degrade the organic d es through the mediation of the h dro l radical under visible light irradiation. More importantl, the CdS quantum dots e hibit good rec cling stabilit during the photocatal tice periments. This suggested that the as-prepared CdS quantum dots might be used as potential photocatal st to effectivel treat the organic pollutants under visible light irradiation.

2012 Elsevier B.V. All rights reserved.

1. Introduction

CdS quantum dots (QDs) have attracted considerable attention over the past decades due to their unique properties and potential applications, such as the photoelectric conversion, light-emitting diodes for flat-panel displas, and lasers [1-3]. However, owing to the ultrasmall si e and high surface energ, the QDs tend to the serious agglomeration, leading to the significant difficulties in the preparation of the stable QDs. Previousl, the CdS QDs were generall prepared using surfactants and pol mers as stabili ers [4-6]. However, the introduction of the to ic stabili ers and the complicated procedures significantle limit the applications. Therefore, it is urgent to construct a facile method to prepare the stable and well dispersed QDs. Because of the coordinating interactions between amino acids and metal ions [7,8], the amino acids might be used as stabiliers to prepare the stable QDs [9–12]. However, hitherto the s nthesis of the stable CdS QDs using h drophobic amino acids as stabili ers is seldom reported and attracts our interest.

Currentl , QDS are widel used in photocatal tic degradation of the biological nondegradable organic pollutants [1–3,13,14]. However, most of the e isting photocatal sts can onl e ert the functions under UV light irradiation, which serious limits their applications. Therefore, it is still a great challenge to find an effective photocatal st to treat the organic pollutants under visible light irradiation.

Herein, using two h drophobic amino acids methionine (Met) and valine (Val) as stabili ers, the stable CdS QDs were successfull prepared through a facile one-pot method. The CdS QDs e hibited special optical properties in visible wavelength region. Moreover, the CdS QDs showed strong photocatal tic degradation activities for organic d es under visible light irradiation. This suggested that the asprepared CdS QDs might be used as photocatal st to effectivel treat the organic pollutants under visible light irradiation.

2. Materials and methods

To prepare the CdS QDs, 0.12 mmol amino acids and 0.06 mmol CdCl $_2$ were dissolved into 20 mL double distilled water (DD water) and incubated for 12 h at 30 °C. Then, 20 mL thioacetamide aqueous solution (3 mM) was added and reflu ed for 1 h at 100 °C. The ellow products were centrifuged, washed, dried in vacuum and denoted as Met-CdS and Val-CdS, respectivel . For comparison, the bulk CdS was also prepared in the absence of amino acids. The si e and morpholog of the samples were determined through transmission electron microscop (TEM, JEOL JEL-2010). The XRD patterns were recorded on DX-2000 diffractometer with Cu K α radiation source ($\lambda = 1.5406\lambda$). The FT-IR spectra were recorded on Bio-Rad FTS-40 infrared spectrometer. The UV–Visible absorption and photoluminescence (PL) spectra were recorded on Lambda-17 UV–vis and Car Eclipse fluorescence spectrophotometers, respectivel .

For photocatal sis e periment, 30 mg CdS QDs were suspended in 24 mL DD water b sonicating. Then 6 mL rhodamine B (RhB) aqueous solution (250 ppm) was added and stirred in the dark for 60 min to establish an adsorption/desorption equilibrium. Then the

^{*} Corresponding authors. Tel.: +86 373 3325058; fa : +86 373 3328507. E-mail addresses: guo uming@gmail.com (Y. Guo), anglin1819@163.com (L. Yang).

suspension was irradiated b 500 W enon lamp for photocatal sis under moderate stirring. At specified intervals, the suspension was taken from the reactor and centrifuged. The content of RhB in the supernatant was determined b UV–Vis spectroscop at 554 nm. For comparison, the photocatal tic activities of the bulk CdS was also determined.

The formation of h dro l radical (•OH) on the photo-irradiated CdS QDs surface was detected *in situ* b fluorimetric assa using terephthalic acid (TPA) as a probe under similar conditions to the photocatal tic e periment e cept the replacement of RhB with TPA. The photo-irradiated suspension was taken from the reactor and centrifuged. Then the fluorescent intensit of 425 nm e cited with 315 nm was detected.

3. Results and discussion

Fig. 1a presents the XRD patterns of the CdS QDs. From the figure, the Met-CdS and Val-CdS e hibit similar XRD patterns, in which three peaks with 20 values of 26.8 , 44.1 and 52.1 correspond to the (1 1 1), (2 2 0) and (3 1 1) planes of the cubic CdS phase (JCPDF 75–0581), respectivel . The broadness of the peaks can be attributed to the

small si es of the QDs. Fig. 1b–e presents the si e and morphologies of the CdS QDs. From the results, the Met-CdS and Val-CdS e hibit similar spherical morpholog but different si es. From the results shown in Fig. 1d–e, the QDs e hibit narrow particle si e distributions. The mean diameter of the Met-CdS is 6.45 nm, smaller than that of the Val-CdS (7.22 nm).

Fig. 2a presents the FT-IR spectra of the bulk CdS and CdS QDs. Compared with the spectrum of the bulk CdS, the stretching bending band of the carbon 1 group of amino acids at 1550–1580 cm⁻¹ can be found in the spectra of the QDs. However, the intensities decrease obviousl . From these results, there is a trace amount of amino acids in the CdS QDs. Furthermore, there are certain interactions between the QDs and the carbon 1 groups of the amino acids. It might be these interactions that ma contribute to the stabilit and dispersit of the CdS QDs.

The optical properties of the CdS QDs, including the UV–visible absorption and PL spectra, were determined and the results are shown in Fig. 2b–c. From Fig. 2b, the CdS QDs e hibit broad absorption band from 200 to 600 nm, indicating the effective photo absorption properties. From the results shown in Fig. 2c, when e cited with wavelength 202 nm, the CdS QDs e hibit a strong green

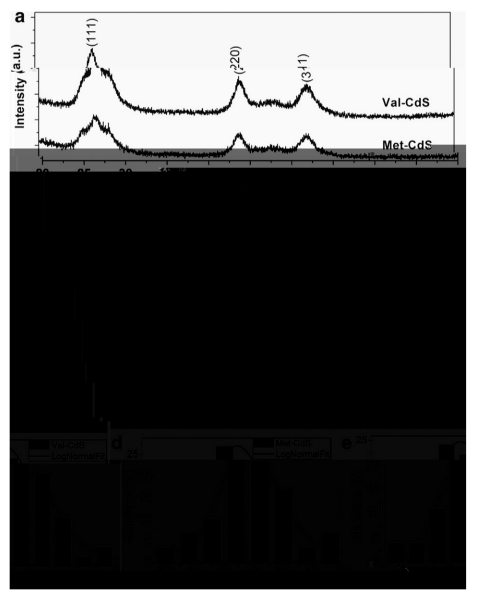


Fig. 1. (a) XRD patterns of the CdS QDs. (b) TEM images of the Met-CdS and (c) Val-CdS QDs. Inset: Magnified images. (d) Particle si e distribution of Met-CdS and (e) Val-CdS QDs.

emission near 528 nm with the quantum ield up to 20%, which can be attributed to the recombination of the charge carriers within surface states.

Because of the photo absorption properties, the photocatal tic ac-

the continuous formation of •OH during the irradiation. Moreover, the PL intensities of the two CdS QDs are different, revealing the different •OH production efficienc . The •OH production efficienc of the ValCdS is higher than that of the Met-CdS, consistent with the photocatal tic activities. Based on the results, the photocatal tic degradation mechanism of the RhB is proposed. Firstl , the photons possessing the energ higher than that of the QDs band gap are firstl absorbed on the QDs surface. This will promote the electrons from the valence band (vb) to the conduction band (cb), generating the positive hole (h_{vb}^+) at the valence band edge and the electron (e_{cb}^-) in the conduction band of the QDs (Eq. (1)). Secondl , h_{vb}^+ and e_{cb}^- can react with water or h dro l groups and transform into •OH (Eqs. (2), (3)). Finall , •OH can react with RhB absorbed on the QDs surface to e ert the degradation (Eq. (4)).

$$CdS + hv \rightarrow CdS(e_{cb}^{-} + h_{vb}^{+})$$
 (1)

$$h_{vb}^{+} + H_2O \rightarrow H^{+} + \bullet OH \tag{2}$$

$$h_{vb}^+ + OH^- \rightarrow \bullet OH$$
 (3)

•OH + dye
$$\rightarrow$$
degradation of the dye (4)

Furthermore, the rec cling of the CdS QDs on the photocatal tic degradation of RhB was determined. From the results (Fig. 3c–d), after five c cles of the photocatal tic reaction, the photocatal tic activities did not show significant loss. These reveal that the CdS QDs can be reused for several times to degrade the organic pollutants under visible light irradiation without great decrease in photocatal tic activit , e hibiting the prospects for practical and long-term applications.

4. Conclusions

In summar , the stable CdS QDs with good optical properties were successfull prepared using Met and Val as the stabili ing agents. Through the mediation of the h dro 1 radicals, the CdS QDs showed strong photocatal tic activities and good rec cling stabilit to degrade the organic d es under visible light irradiation, suggesting the potential application in the effectivel treatment of the organic pollutants under visible light irradiation.

Acknowledgements

This work was financiall supported b the National Science Foundation of China (21171051) and the Program for Changjiang Scholars and Innovative Research Team in Universit (IRT1061) and the Innovation Fund for Outstanding Scholar of Henan Province (114200510004) and Henan Ke Proposed Program for Basic and Frontier Research (112102210005, 112300410095) and the Natural Science Foundation of Henan Educational Committee (2010A150013) and the Henan Normal Universit College Students Innovation E periment Program.

References

- [1] Fan Y, Deng M, Chen G, Zhang Q, Luo Y, Li D, et al. J Allo s Compd 2011;509: 1477–81.
- [2] Hu Y, Liu Y, Qian H, Li Z, Chen J. Langmuir 2010;26:18570-5.
- [3] Rajendran V, K nig A, Rabe KS, Nieme er CM. Small 2010;6:2035–40.
- [4] Ente ari MH, Ghows N. Ultrason Sonochem 2011;18:127–34.
- [5] Ma N, Doole CJ, Kelle SO, J Am Chem Soc 2006;128:12598-9.
- [6] Yoon H, Lee J, Park D, Hong C, Shim S. Colloid Pol m Sci 2010;288:613-9.
- [7] Chin J, Lee SS, Lee KJ, Park S, Kim DH. Nature 1999;401:254–7.
- [8] Davis AV, O Halloran TV. Nat Chem Biol 2008;4:148–51.
- [9] Cao X, Li CM, Bao H, Bao Q, Dong H. Chem Mater 2007;19:3773-9.
- [10] Chanu TI, Negi DPS. Chem Ph s Lett 2010;491:75–9.
- [11] Guo Y, Wang L, Yang L, Zhang J, Jiang L, Ma X. Mater Lett 2011;65:486-9.
- [12] Park Y-S, Dm truk A, Dmitruk I, Kasu a A, Takeda M, Ohuchi N, et al. ACS Nano 2009;4:121–8.
- [13] Guo Y, Wang J, Yang L, Zhang J, Jiang K, Li W, et al. Cr stEngComm 2011;13: 5045–8.
- [14] Wang J, Guo Y, Yang L, Zhang J, Chen D. Curr Nanosci 2010;6:539-44.