Synthesis, Optical Characterization, and Size Distribution Determination by Curve Resolution Methods of Water-Soluble CdSe Quantum Dots

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In this work a colloidal approach to synthesize water-soluble CdSe quantum dots (QDs) bearing a surface ligand, such as thioglycolic acid (TGA), 3-mercaptopropionic acid (MPA), glutathione (GSH), or thioglycerol (TGH) was applied. The synthesized material was characterized by X-ray diffraction (XRD), Fourier-transform infrared spectroscopy (FT-IR), UV–visible spectroscopy (UV-Vis), and fluorescence spectroscopy (PL). Additionally, a comparative study of the optical properties of different CdSe QDs was performed, demonstrating how the surface ligand affected crystal growth. The particles sizes were calculated from a polynomial function that correlates the particle size with the maximum fluorescence position. Curve resolution methods (EFA and MCR-ALS) were employed to decompose a series of fluorescence spectra to investigate the CdSe QDs size distribution and determine the number of fraction with different particle size. The results for the MPA-capped CdSe sample showed only two main fraction with different particle sizes with maximum emission at 642 and 686 nm. The calculated diameters from these maximum emission were, respectively, 2.74 and 3.05 nm.

Keywords: Semiconductors, Optical materials, CdSe, Optical properties, size distribution

1. Introduction

Colloidal semiconducting nanocrystals, also known as quantum dots (QDs), have attracted widespread attention over the last decade-QDs display novel optical properties such as intense luminescence, narrow and symmetric emission, broad and strong absorption, and high stability in presence of photo and chemical radiation.1 QDs are particularly interesting for fundamental studies in the field of nanoscience, have potential technological applications in optoelectronic devices, including light emitting diodes (LEDs), lasers, photodetectors, and photovoltaic cells,2,3 and may aid labeling of biomedical entities.6 Quantum confinement effects give rise to semiconducting nanoparticles with physical and chemical properties that are distinct from the properties of the individual molecule or the bulk species.7 The possibility of confining charge carriers (electrons and holes) in three dimensions endows these intrinsic semiconductors with a size-dependent band gap and with optical properties that strongly rely on nanoparticle size and structure.8,9 Kamat et al.10 reported on the specific advantages of using QDs as light harvesting assemblies in solar cells. The size effect allowed these authors to tune the visible response, vary the band offsets, and modulate the vectorial charge transfer across particles of different sizes. This effect could pave the way for using hot electrons or generating multiple charge carriers with a single photon.

Cadmium selenide (CdSe) is one of the most important materials among the various kinds of II-VI semiconductors. Its size-tunable emission in the visible range has made it a useful model system to investigate a wide array of electronic, optoelectronic, and chemical processes at nanoscale.11 CdSe QDs display a direct band gap of 1.74 eV, which makes their use in solar cells and other electronic devices feasible.10-14 Reactions in organic media constitute the main route to obtain CdSe QDs.15-17 These synthetic approaches produce high-quality optical nanoparticles18 with narrow size distribution.15,19 The preparation of CdSe QDs in aqueous media is reproducible and inexpensive, and the resulting water-soluble QDs present low toxicity in comparison with that produced in organic media.11,20,21 Because the large absorption and emission bands of water-soluble CdSe nanocrystals remain poorly understood, there is little information about their size distribution. Exchange of the short-chain ligand for a long-chain ligand renders water-soluble CdSe QDs easily soluble in organic solvents,22 as shown by Gégout et al.23 Obtaining nanocrystals that are compatible with organic solvents is vital for applications in optoelectronic devices.

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All these advantages have motivated research into water-soluble QDs in an attempt to increase the photoluminescence quantum yield ($\Phi_L$) of materials synthesized in aqueous medium and to promote surface modifications that will ensure desirable optical and electronic properties as well as good chemical stability.

Regarding the synthesis of QDs via the water-based method, it is possible to alter the size-dependent features and structure of the nanocrystals by modifying a set of variables such as precursors concentration, reaction time, reaction medium pH, reaction temperature, surface ligand type, and reactants molar ratio, among other parameters.  

Many methods such as high-resolution transmission electron microscopy (TEM), 24 small-angle X-ray scattering, 25 Raman scattering, 26 and UV-vis absorption spectroscopy 29 can help to determine the accurate size distribution of QDs. Unfortunately, these methods still have limitations like time-consuming sample preparation, limited instrument availability, laborious data processing, and low resolution. 27 The color of the QD emission depends on nanocrystal size, while the width of the PL peak depends on nanocrystal size distribution. Hence, on the basis of previous studies, correlation fluorescence spectra could afford the maximum diameter of QDs. 28

Peng et al. 26 correlated the optical properties of QDs with nanocrystal size. In the case of high-quality CdTe, CdSe, and CdS nanocrystals, these authors found that the extinction coefficient per mole of nanocrystals at the first excitonic absorption peak heavily depended on crystal size (between a square and a cubic dependence). By compiling TEM data and the maximum of the first excitonic absorption peak, these authors established a useful empirical dependence that aided determination of the size and extinction coefficients of QDs. Mulvaney et al. 29 re-examined the size-dependent optical absorption coefficients of CdSe nanocrystals at the band-edge and constructed calibration curves for the size-dependent first absorption peak and the molar extinction coefficient. These calibration curves helped to determine the concentration of nanocrystals in solution accurately.  

Radotic et al. 27 developed the first method to estimate the size of QDs on the basis of their emission properties. These authors employed statistical methods (EFA-evolving factor analysis and MCR-ALS-multivariate curve resolution-alternating least square) to decompose the emission spectra of nanoparticles and identify fractions of QDs with different diameters. Recent works have highlighted this methodology. 28,30

Together, MCR-ALS 31-33 and EFA 34-35 are some of the most popular methods to deconvolution overlapping signals.  

MCR-ALS can handle multiple arrays simultaneously, thereby reducing the rotational ambiguity intrinsic to bilinear methods and eliminating problems of rank deficiency. Moreover, MCR-ALS presents diverse and flexible constraints that help to improve resolution. 31 Finally, MCR-ALS has a very friendly graphical user interface 37 with easy and free access through the MATLAB platform.

EFA is a time dependent rank analysis where the evolution of the total rank of a matrix X is followed by plotting the singular values of an increasing number of consecutive spectra as a function time. 34,35 This approach allows determination of the number of components, which here is related to the number of fraction of QDs with different distribution sizes, and gives an initial estimate of concentration profile.  

In this perspective, this study deals with the synthesis, optical characterization and size distribution determined indirectly by curve resolution methods EFA and MCR-ALS applied to photoluminescence data of CdSe QDs in colloidal aqueous medium using different reaction times and different surface ligands.

2. Experimental

2.1. Materials

Selenium powder (100 mesh, 99.8%), thioglycolic acid (TGA, 99%), 3-mercaptopropionic acid (MPA, 99%), glutathione (GSH, 99%) and thioglycerol (TGH, 98%) were obtained from Sigma-Aldrich (St. Louis, MO, USA). Sodium borohydride (NaBH₄, 97%) was acquired from Nuclear (São Paulo, Brazil). Cadmium chloride monohydrate (CdCl₂·H₂O, 99%) was provided by Vetec (Rio de Janeiro, Brazil). All the chemicals were used without additional purification. Milli-Q ultrapure water (Millipore Co., Billerica, MA, USA) was employed in the synthesis of nanoparticles.

2.2. Synthesis of CdSe QDs

The experimental procedure was performed on the basis of previous studies. 39,41,42,43 First, the selenium precursor was prepared in a three neck round-bottom flask by dissolving 0.4 mmol of selenium and 1 mmol of sodium borohydride in 10 mL of distilled water, under argon atmosphere and constant stirring. This solution was kept in an ice bath for 2 h, until all the selenium was reduced. In a second reaction flask, 0.4 mmol of cadmium chloride and 2 mmol of ligand was added to 80 mL of distilled water; the pH was adjusted to 10.0 by addition of sodium hydroxide. Next, the selenium precursor solution, as prepared, was added to the latter reaction flask, and the resulting solution was stirred and heated (100 °C). Particle size was monitored throughout the process. To follow the progress of the synthesis, aliquots were removed from the reaction medium at regular time intervals and submitted to UV-Vis and PL analyses. Four different types of surface ligands were used: 3-mercaptopropionic acid (MPA), thioglycolic acid (TGA), thioglycerol (TGH) and glutathione (GSH) (Figure 1).

At the end of the heating and stirring process, the CdSe solution was separated from the reaction mixture. First, the as-prepared nanocrystal solution was concentrated to one fifth of its volume in a rotary evaporator attached...
to a vacuum pump. Then, a small portion of acetone was added to the solution, to remove impurities and other reaction products. The supernatant of this solution was separated by centrifugation, and acetone was added to the supernatant again, until the solution became cloudy. Further centrifugation allowed larger CdSe particles to precipitate. The precipitated particles were dried at 40°C, in an oven.

2.3. Characterization

UV-Vis absorption and fluorescence at single excitation wavelength spectra were acquired on a Varian Cary 50 spectrophotometer and on a Shimadzu RF-5301 PC spectrofluorophotometer, respectively. Absorption and fluorescence measurements were performed in quartz cuvettes with an optical path length of 10 mm (Shimadzu). Spectra of air-saturated solutions were recorded at room temperature. X-ray diffraction (XRD) patterns of the CdSe QD samples sprayed in acetone were registered in a non-reflective silicon sample port with the aid of a Shimadzu XRD-6000 diffractometer operating in the scanning mode (15º-60º) and using CuKα radiation (λ = 1.5418 Å) generated at 40 kV and at a current of 30 mA. Steps of 0.02° and a sampling time of 4 h were employed. Divergence slits of 1.0 mm, scattering slits of 1.0 mm, and receiving slits of 0.3 mm were used. Fourier-transform infrared spectroscopy (FT-IR) was conducted on a spectrometer Spectrum GX Perkin Elmer. The spectra were obtained in the transmission mode in the range of 400 to 4000 cm⁻¹ with accumulation of 32 spectra. The resolution was 2 cm⁻¹. Additionally, the multi-wavelength fluorescence data were obtained from the successive acquisition of emission spectra at multiple excitation wavelengths, generating excitation-emission matrices (EEM) for CdSe QDs bearing distinct surface ligands. For CdSe/MPA QDs, the EEM were collected for each time of synthesis with spectral ranges of 440-540 nm for excitation and 550-900 nm for emission. For CdSe/GSH, the spectral range were 410-510 nm for excitation and 400-900 nm for emission.

2.4. Size distribution determination of CdSe QDs

The prepared CdSe QDs were re-dispersed in water prior to the absorption and fluorescence emission experiments. First, the CdSe QD sizes were determined by UV-Vis spectroscopy with the aid of the equations obtained by Peng et al.°. To this end, samples were transferred to quartz cuvettes, and the absorption spectrum was recorded. Calibration curves were used to correlate the size of CdSe QDs with the maximum fluorescence.

The successive acquisition of emission spectra at multiple excitation wavelengths, which generates excitation-emission matrices (EEM), provides full spectral information and is more accurate than absorption spectra. In this sense, specific measurement procedures were applied to detect any signal of fluorescence for CdSe QDs bearing distinct surface ligands.

On average, emission spectra were collected for each CdSe QDs dispersion obtained at different synthesis time of reaction by varying the excitation wavelength by 5 nm. To include all the geometric characteristics of the emission spectrum, for each sample, the initial excitation wavelength was defined in such a way that recording of the emission spectra began between 50 and 60 nm before the emission increased sharply.

In order, to get a complete spectral information of each CdSe QDs dispersion, excitation-emission matrices were also collected at different times of the synthesis -15, 30, 60, 90, 120, 150, and 180 min. For the matrices whose excitation and emission wavelengths were overlapped, it was necessary to accomplish a pre-treatment that eliminated the elastic Rayleigh scattering. This pre-treatment excluded the affected region and recreated this area by interpolation. Then, the matrices were arranged into single ones by matrix column-augmentation (X = [X15; X30; X60; X90; X120; X150; X180]), and curve resolution methods were applied to investigate the CdSe QDs size distribution and determine the number of fraction with different particle size.

MCR-ALS is an iterative curve resolution method that focuses on the bilinear decomposition of mixed multicomponent information store in a data matrix into chemically meaningful pure contributions of each of the components in the mixture. The bilinear decomposition of the experimental data matrix is performed using the model equation:

$$X = CS^T + E,$$

where $C$ is the matrix that describes how the excitation spectra of the $n$ QDs changes along the time, and $S^T$ is the matrix that represents the pure emission spectra of the $n$ QDs of the data matrix. $E$ is the residual matrix corresponding to the variance not explained by the model $CS^T$.

The first step in the resolution procedure was to determine the number of components, $n$, causing the observed variance in $X$. The second step was to obtain an initial estimate.
of the concentration matrix. In this work, the number of components, \( n \), and, the initial estimate of concentration matrix were obtained by the EFA method. The third step referred to the MCR-ALS decomposition step. At this stage, the restriction of non-negativity of the spectral profiles was implemented. The MCR-ALS program implemented in the MATLAB 7.0 platform was used for all calculations.

3. Results and Discussion

3.1. CdSe QDs

A colloidal approach that used distinct surface ligands afforded different CdSe QDs in aqueous medium. A comparative study of these CdSe QDs showed how the surface ligands TGA, MPA, GSH, and TGH affected the optical properties of the nanocrystals.

To obtain the CdSe QDs, the initial step was to react metallic selenium and sodium borohydride, to achieve the NaHSe precursor as represented by reaction (1): 

\[
4\text{NaBH}_4 + 2\text{Se} + 7\text{H}_2\text{O} \rightarrow 2\text{NaHSe} + \text{Na}_2\text{B}_4\text{O}_7 + 14\text{H}_2
\]  

(1)

Then, the NaHSe precursor was added to a solution containing the surface ligand and Cd\(^{2+}\) ions, to give the CdSe nanoparticles according to reaction (2): 

\[
\text{NaHSe} + \text{CdCl}_2 \rightarrow \text{CdSe} + \text{Na}^+ + 2\text{Cl}^- + \text{H}^+
\]  

(2)

Figure 2 depicts the UV-vis and fluorescence spectra that helped to monitor the progress of the synthesis of CdSe QDs bearing MPA as surface ligand (UV-vis and fluorescence spectra of CdSe QDs synthesized using GSH, TGA e TGH as surface ligands are shown in Figures 3, 4 and 5). For all the samples, the excitonic absorption and luminescence bands shifted slightly toward lower energies with increasing particle size. This shift was consistent with quantum confinement effects and agreed with literature results published for similar systems. Figure 2a revealed slightly red-shifted absorption bands on going from the aliquot withdrawn at 15 min of synthesis, 482 nm, to the aliquot withdrawn at 180 min of synthesis, 493 nm. This shift suggested that nanoparticle size increased along the synthesis. Similarly, the PL spectra, obtained at excitation wavelength of 355 nm, in Figure 2b showed slight displacement of the emission bands toward longer wavelengths, from 649 nm in the case of the first aliquot to 659 nm at the end of the synthesis. This displacement corresponded to the emission of orange color.

Figure 6 illustrates the temporal evolution of the position of the absorption and emission bands for the different surface ligands used during the growth of CdSe QDs. GSH shifted the absorption (\(~ 53\) nm) and emission (\(~ 140\) nm) bands toward the red the most. In other words, this capping agent provided higher nanoparticle growth rate as compared with the other capping agents. This faster growth may have originated from the higher steric hindrance and the larger susceptibility to thermal decomposition of GSH as compared with the other ligands. Hence, GSH was more turned off from the surface of the QDs (“off” state). Conversely, TGH displaced the absorption (~ 10 nm) and emission (~ 6 nm) bands the least, which indicated that this ligand promoted the slowest kinetic growth of CdSe nanocrystals. Probably, the two close functional OH groups in the structure of TGH caused this ligand to be more strongly attached to the surface of the nanocrystals (“on” state), thereby preventing a large increase in particle size. As for MPA and TGA, results were similar and lay between the behaviors of GSH and TGH, as seen from the shifts in the absorption (~ 20 nm and ~ 29 nm for MPA and TGA, respectively) and emission (~ 12 nm and ~ 13 nm for MPA and TGA, respectively) bands. For TGA-capped QDs a NIR (near infra-red) emission could be related to the surface-defects that generate shallow levels near the valence band, that acts as traps for photogenerated holes. The obtained results agreed with literature data presented for CdTe QDs obtained in aqueous medium and bearing GSH, MPA, TGA, and TGH as surface ligands.

Figure 7 brings the XRD patterns registered for the CdSe QDs stabilized with the studied ligands. The diffraction patterns displayed the three main diffraction peaks of CdSe, at 2\(\theta\) = 25.1, 41.8, and 49.2 degrees, which confirmed the crystallinity of the CdSe QDs. The positions of the diffraction peaks standing for the (111), (220), and (311) planes referred to the zinc blend structure and matched the standard pattern well (JCPDS No.– 00-019-0191). Among the employed surface ligands, GSH shifted the diffraction peaks toward the CdS cubic crystalline phase the most. This happened because GSH was more susceptible to thermal decomposition and probably passivated the QDs surface more effectively. This may have increased sulfur release on the surface of the CdSe nanocrystals and may even have generated a thin CdS layer, to give a quasi core-shell structure. According to the literature, CdTe QDs capped with GSH in aqueous medium display the same behavior.

Infrared spectroscopy furnished information about the bond that originated between the surface ligands and the CdSe surface. For illustration, Figure 8 contains the infrared spectra of the pure MPA, TGA, TGH, and GSH ligands and of the CdSe QDs stabilized with these ligands. Additionally, Figure 8 presents the assignment of the main FT-IR band and, holes. near the valence band, that acts as traps for photogenerated holes.
due to C–O stretching. In contrast, the spectra of the capped CdSe samples displayed the symmetric and asymmetric absorption bands assigned to the carboxylate anion of the surface ligands. The existence of the carboxylate anion in the QDs resulted from the synthesis process, which used NaOH to adjust the pH to 10.0 and therefore deprotonated the COOH group.

The O–H stretching of absorbed water emerged as a broad band in the region of 3300–2900 cm$^{-1}$ in the spectra of MPA-capped CdSe QDs, TGA-capped CdSe QDs, and TGH-capped CdSe QDs, whereas the spectrum of GSH-capped CdSe QDs exhibited a broad band in the region of 3500–3100 cm$^{-1}$, attributed to N–H stretching. The spectrum of TGH-capped CdSe QDs also had absorption

Figure 2: UV-visible absorption (a) and fluorescence (b) spectra of CdSe/MPA QDs at different reaction times.

Figure 3: UV-visible absorption a) and fluorescence b) spectra of CdSe/GSH QDs at different reaction times.

Figure 4: UV-visible absorption a) and fluorescence b) spectra of CdSe/TGA QDs at different reaction times.
bands around 1500 and 900 cm\(^{-1}\), ascribed to O–H bending and C–O stretching.\(^{50,51}\) The FT-IR assignments agreed with data recorded for similar systems such as water-soluble CdTe, which means that CdSe and CdTe QDs interact with thiol surface ligands in aqueous medium in much the same way.

### 3.2. Size determination of CdSe QDs

The absorption wavelength (\(\lambda_{\text{abs}}\)) obtained by UV-Vis spectroscopy and the equation obtained by Peng et al.\(^{26}\) aided size determination of CdSe QDs. Table 1 shows the correlation between the calculated diameters and the maximum emission wavelength for MPA-capped CdSe QDs, at excitation wavelength of 355 nm.

The data presented in Table 1 allowed construction of a calibration curve that related the position of the maximum fluorescence with the diameter calculated for the nanoparticle (see Figure 9 for the calibration curves constructed for MPA-capped CdSe QDs and GSH-capped CdSe QDs). A polynomial fit of the experimental data provided a better calibration curve, yielded the highest coefficient of determination\(^{28}\) (\(R^2\)) and the deviation from the data, and gave a fine description of the system in the treated interval. In this way, by the calibration curve equation, and knowing the sample maximum fluorescence position, is possible to estimate the average particles sizes.

Fluorescence was measured for each capped CdSe QDs sample at different times; i.e., at 15, 30, 60, 90, 120, 150, and 180 min of synthesis. Because QDs of different sizes should
Synthesis, Optical Characterization, and Size Distribution Determination by Curve Resolution Methods of Water-Soluble CdSe Quantum Dots

Figure 8: Infrared spectra of CdSe QDs capped with (a) MPA, (b) TGA, (c) TGH, and (d) GSH.

Table 1: Position of the maximum fluorescence in relation to the diameter of MPA-capped CdSe QDs.

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<tr>
<th>Time</th>
<th>Maximum Fluorescence position (nm)</th>
<th>Particle diameter (nm)</th>
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<tr>
<td>15 min</td>
<td>648</td>
<td>2.07</td>
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<tr>
<td>30 min</td>
<td>652</td>
<td>2.16</td>
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<tr>
<td>180 min</td>
<td>660</td>
<td>2.23</td>
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emit in different positions of the spectrum, this procedure helped to determine the size distributions of the QDs along the synthesis. In addition, the samples were irradiated with different wavelengths by varying the excitation wavelength by 5 nm each time the spectrum was recorded. Figure 10 illustrates the results for 30 min MPA-capped CdSe QDs and 30 min GSH-capped CdSe QDs.

Decomposition by the MCR-ALS method of the set of spectra obtained for MPA-capped CdSe QDs revealed the existence of two types of QDs with specific diameters in this material (Figure 11a). More specifically, MPA-capped CdSe QDs displayed maximum emission at 642 and 686 nm, which corresponded to calculated diameters of 2.74 ± 0.1 and 3.05 ± 0.1 nm, the calculated average particles sizes, obtained after the distribution of sizes calculations, presents an error of 0.1 nm, due to the calibration curve equation error, as inferred from the plot depicted in Figure 9. As for GSH-capped CdSe QDs, maximum emission emerged at 582 and 658 nm, which corresponded to diameters of 1.94 ± 0.1 and 2.32 ± 0.1 nm, respectively (Figure 11b). With respect to the synthesis method, and synthesis time, GSH-capped CdSe QDs and MPA-capped CdSe QDs showed the same size distribution with two resolved emission spectra for each system. This is an important result: while the very large bandwidth values of water-soluble CdSe are mainly associated with broad size distribution, here there were only two main sizes of capped CdSe QDs at different times of synthesis. Therefore, the structure and shape of the nanocrystals in the CdSe QDs samples synthesized in this work were similar. Actually, the different thiol-based surface ligands affected the nanocrystal growth rate, but they did not influence the nanocrystal size distribution.
4. Conclusions

This work described the successful synthesis of capped CdSe QDs via a chemical colloidal process conducted in aqueous medium, in the presence of one of the following surface ligands: thioglycolic acid (TGA), 3-mercaptopropionic acid (MPA), glutathione (GSH), or thioglycerol (TGH). A comparative study showed that the MPA- and TGA-capped CdSe QDs displayed similar properties. As for the TGH ligand, it bears two functional OH groups in close proximity in its structure. These groups promoted strong interactions between TGH and the nanocrystals surface, preventing CdSe growth. The size of the GSH ligand provided faster CdSe QDs growth. In addition, the thermal instability of GSH during the synthesis of capped CdSe QDs released sulfur on the...
surface of the nanocrystals and generated a quasi CdSe/CdS core/shell structure. The capping agent probably attached to the CdSe QDs surface via the –SH end of the molecule.

Decomposition of the spectra of the various CdSe QDs by chemometric techniques afforded the size distribution of the nanoparticles obtained during the synthesis. GSH-capped CdSe QDs and MPA-capped CdSe QDs showed the same size distribution, with two resolved emission spectra for each surface ligand. The maximum emission bands obtained for GSH-capped CdSe QDs were 582 and 658 nm, which corresponded to QDs with diameters of 1.94 and 2.32 nm, respectively. As for MPA-capped CdSe QDs, the maximum emission bands were 642 and 686 nm, referring to QDs with diameters of 2.74 and 3.05 nm, respectively.

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6. References


