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Ex situ formation of metal selenide quantum dots using bacterially derived selenide precursors

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Abstract
Luminescent quantum dots were synthesized using bacterially derived selenide (SeII−) as the precursor. Biogenic SeII− was produced by the reduction of SeIV by Veillonella atypica and compared directly against borohydride-reduced SeIV for the production of glutathione-stabilized CdSe and β-mercaptoethanol-stabilized ZnSe nanoparticles by aqueous synthesis. Biological SeII− formed smaller, narrower size distributed QDs under the same conditions. The growth kinetics of biologically sourced CdSe phases were slower. The proteins isolated from filter sterilized biogenic SeII− included a methylmalonyl-CoA decarboxylase previously characterized in the closely related Veillonella parvula. XAS analysis of the glutathione-capped CdSe at the S K-edge suggested that sulfur from the glutathione was structurally incorporated within the CdSe. A novel synchrotron based XAS technique was also developed to follow the nucleation of biological and inorganic selenide phases, and showed that biogenic SeII− is more stable and more resistant to beam-induced oxidative damage than its inorganic counterpart. The bacterial production of quantum dot precursors offers an alternative, ‘green’ synthesis technique that negates the requirement of expensive, toxic chemicals and suggests a possible link to the exploitation of selenium contaminated waste streams.

1. Introduction

The use of microorganisms for the synthesis of nanocrystalline particles represents a low cost, environmentally-friendly method of producing industrially and technologically relevant nanomaterials. Microorganisms have been used to successfully synthesize a range of nanomaterials including Fe oxides, precious metal catalysts and a suite of nanocrystalline, semiconducting ‘quantum dots’ [1–4].

The interest in quantum dots (QDs) is due to their unique semiconducting properties induced by the onset of quantum confinement in the size range 1–20 nm, generating optical, electrical and mechanical properties that differ from those of the bulk materials [5]. The photooptical and photovoltaic properties of the II/VI semiconductors (metal chalcogenides) are particularly suited for their application in solar cells and optoelectronic sensors [5–7] as well as fluorescent biolabelling, including their use in cancer screening [8–10].
Of all QDs synthesized, cadmium-based QDs (CdS, CdSe and CdTe) are the most thoroughly explored owing to fluorescent emissions across the visible spectrum by direct manipulation of particle size.

Traditional organochemical synthesis of chalcogenide quantum dots involves the injection of reactants into heated organic solvents, commonly tri-n-octylphosphine oxide (TOPO). Synthesis requires the use of a number of toxic, expensive precursors and necessitates the exchange of surfactant molecules to aid aqueous solubility [11, 12]. Simpler, aqueous-based synthesis techniques have subsequently been developed in which quantum dots are produced in a reproducible, single-step reaction at temperatures <100 °C, using surfactant molecules to increase water solubility and stability, and to limit particle growth [7, 11, 13]. Thiol compounds have been identified as ideal surfactants, and investigations by Rogach et al [14] and Gaponik et al [7] identify thioalcohols such as β-mercaptoethanol as especially suited to minimizing the size of initial precipitates. Biologically derived sulfurous materials such as the peptides glutathione (GSH) and bovine serum albumin (BSA) have also been used successfully in the aqueous synthesis of metal chalcogenide quantum dots [11, 15–21].

The whole-cell biological synthesis of II/VI quantum dots has been reported as a greener, environmentally-friendly alternative to chemical synthesis. The use of a number of fungal [22–26] and bacterial [18, 27–30] species in the formation of a range of quantum dots has been demonstrated (table 1), and exploit potential heavy metal detoxification mechanisms. CdII detoxification by yeast species including Schizosaccharomyces pombe, Candida glabrata [15, 16] and Saccharomyces cerevisiae [26] stimulates the secretion of extracellular sulfurous proteins which act as surfactants, as well as stimulating sulfide generation, resulting in extracellular deposition of peptide-coated fluorescent CdS nanoparticles. The use of oxanions of the chalcogens Se and Te as precursors typically requires the addition of a strong reducing agent such as sodium borohydride to produce the required SeII or TeII ions [1, 19, 21, 26, 28, 31], however, Pearce et al [32] demonstrated the ability of the clinical isolate Veillonella atypica to reduce aqueous SeIV to SeII through a biphasic reduction pathway, suggesting the possibility of linking the biosynthesis of quantum dot precursors to the bioremediation of selenium contaminated waste streams [33]. Further research on this bacterium demonstrated the feasibility of the use of biologically prepared aqueous SeII– solutions in the formation of metal selenide quantum dots [18].

This research reports on the potential for tailoring the ex situ formation of metal selenide quantum dots using bacterially generated SeII– as an alternative to chemically synthesized precursors. Biogenic and abiotic fluorescent nanoparticles were characterized and compared using UV/visible light (UV/vis) spectrophotometry, photoluminescence (PL) spectroscopy and transmission electron microscopy (TEM). Bacterial proteins present within filter sterilized biogenic SeII– solutions were identified by polyacrylamide gel electrophoresis. Potential chemical interactions between the nanoparticles and glutathione was explored using S K-edge synchrotron x-ray absorption spectroscopy (XAS). Finally, this work describes the effects of extracellular biological materials on the stability, nucleation and growth of β-mercaptoethanol stabilized, zinc selenide nanoparticles using a novel in situ time-resolved synchrotron XAS technique.

2. Methods

All chemicals used in this work were of analytical grade and acquired from Sigma-Aldrich, UK.

2.1. Bacterial synthesis of SeII–

Veillonella atypica (ATCC 14894) was grown anaerobically in defined media, coupling the reduction of 1.0 g l–1 glucose to the oxidation of 7.5 g l–1 sodium lactate [18]. Cells were isolated by centrifugation and re-suspended in sterile, anaerobic 20 mM 3-(N-morpholino)propanesulfonic acid (MOPS) buffer at pH7.5 to an OD600 of ~1.0. Resting cell cultures were amended with 5 mM Na2SeO3 and 100 µM of the electron shuttling compound anthraquinone-2,6-disulfonate (AQDS). The head spaces of the sealed culture bottles were identified as ideal surfactants, and investigations by Rogach et al [14] and Gaponik et al [7] identify thioalcohols such as β-mercaptoethanol as especially suited to minimizing the size of initial precipitates. Biologically derived sulfurous materials such as the peptides glutathione (GSH) and bovine serum albumin (BSA) have also been used successfully in the aqueous synthesis of metal chalcogenide quantum dots [11, 15–21].

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Table 1. A number of fungal and bacterial species have been used in the synthesis of a number of metal chalcogenide nanoparticles.

<table>
<thead>
<tr>
<th>Kingdom</th>
<th>Microorganism</th>
<th>Product</th>
<th>Metal reactant</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fungi</td>
<td>Fusarium oxysporum</td>
<td>CdSe</td>
<td>CdCl2</td>
<td>[23]</td>
</tr>
<tr>
<td></td>
<td>Candida glabrata</td>
<td>CdS</td>
<td>CdSO4</td>
<td>[22]</td>
</tr>
<tr>
<td></td>
<td>Schizosaccharomyces pombe</td>
<td>CdS</td>
<td>CdSO4</td>
<td>[22, 24]</td>
</tr>
<tr>
<td></td>
<td>Torulopsis sp.</td>
<td>PbS</td>
<td>Pb(NO3)2</td>
<td>[25]</td>
</tr>
<tr>
<td></td>
<td>Saccharomyces cerevisiae</td>
<td>CdTe</td>
<td>CdCl2</td>
<td>[26]</td>
</tr>
<tr>
<td></td>
<td>Escherichia coli</td>
<td>CdS</td>
<td>CdCl2</td>
<td>[30]</td>
</tr>
<tr>
<td></td>
<td>Klebsiella pneumoniae</td>
<td>CdS</td>
<td>Cd(NO3)2</td>
<td>[29]</td>
</tr>
<tr>
<td>Bacteria</td>
<td>Escherichia coli</td>
<td>CdTe</td>
<td>CdCl2</td>
<td>[28]</td>
</tr>
<tr>
<td></td>
<td>Rhodobacter sphaeroides</td>
<td>ZnS</td>
<td>ZnSO4</td>
<td>[27]</td>
</tr>
<tr>
<td></td>
<td>Veillonella atypica</td>
<td>CdSe</td>
<td>Cd(ClO4)2</td>
<td>[18]</td>
</tr>
</tbody>
</table>
0.22 \mu m micropore filters. The filtered biogenic Se^{II-} solutions were then split into 15 ml aliquots and the pH was adjusted to 11.2 with NaOH.

2.2. Abiotic synthesis of Se^{II-}

Abiotic sodium hydrogen selenide was synthesized by adding sodium borohydride (1.15 g, 30.5 mM) in 12.5 ml of degassed deionized water to grey selenium powder (1.15 g, 14.5 mM) suspended in 12.5 ml degassed deionized water under an \textsubscript{N}_2 atmosphere, at room temperature and with stirring [34]. The resulting, virtually colourless solution of NaHSe was filtered to remove crystals of Na_2B_4O_7·10H_2O that formed upon cooling of the solution after the exothermic reaction.

The NaHSe stock solution was diluted to 5 mM with 20 mM MOPS buffer, and the pH was raised to 11.2 using NaOH under an \textsubscript{N}_2 atmosphere.

2.3. Formation and growth of CdSe quantum dots

A solution of 10 mM Cd(ClO\textsubscript{4})\textsubscript{2} and 30 mM reduced glutathione (GSH) was prepared so that the final molar ratio upon mixing would be 2:1:3 (Cd:Se:GSH). High purity N\textsubscript{2} was bubbled through the Cd^{II+}-GSH solution to remove O\textsubscript{2}. The pH was then adjusted using NaOH under an \textsubscript{N}_2 atmosphere to 11.2. Solutions of abiotic and biogenic Se^{II-} and Cd^{II+}-GSH were chilled in an ice bath for 1 h. Aliquots of the Se^{II-} solutions were injected rapidly into equal volumes of the Cd^{II+}-GSH solutions under an oxygen-free atmosphere, and were shaken vigorously for 30 s until a yellow precipitate was observed. The GSH–CdSe suspension was then transferred in to a three-necked round bottom glass flask, and refluxed on a heating mantle at 100\degree C. Aliquots were taken at 4 min time intervals until no suspension remained. Aliquots were immediately quenched in liquid nitrogen and stored at −80\degree C prior to analysis.

2.4. Characterization of CdSe quantum dots

The UV/vis absorption spectra of the reaction suspension aliquots were measured with an Analytik Jena Spectrocolor S600 (Analytik Jena AG, Germany). Photoluminescence spectra were taken with a Gilden Photonics Fluorosens (Guilden Photonics Ltd, UK), using a constant excitation wavelength of 365 nm.

HR-TEM images were taken with an FEI Tecnai F20 FEIGTEM (200 kV), fitted with an Oxford Instruments 80 mm\textsuperscript{2} XMAX SDD EDX detector and a Gatan Orius SC200 CCD. TEM samples were prepared by dropping an aliquot of QDs in water onto a 400-mesh carbon-coated copper grid (Agar Scientific, UK). Particle sizing was via analysis of TEM images, with more than 100 particles assessed at each sample time point for statistical accuracy.

2.5. Protein identification

To analyse for protein in the selenide solutions, 25 \mu l fractions of the abiotic and biogenic Se^{II-} solutions were run in an 18% Tris–HCl gel with Tris/glycine/SDS running buffer. Kaleidoscope (BioRad) prestained molecular weight markers were run on both sides of the gel and gel-separated protein bands were detected using Imperial Protein Stain. Bands of interest were excised under sterile conditions for trypsin digestion. Tryptic peptides were analysed using liquid chromatography-based high-resolution tandem mass spectrometry (LC-MS/MS), comprising an Ultimate 3000 (LC-Packings, Dionex, Amsterdam, The Netherlands) coupled to a HCT Ultra ion trap mass spectrometer (Bruker Daltonics, Bremen, Germany). All samples were searched first against the SWISSPROT database searching all species, and then against either a species-specific search in UNIPROT, or against a species-specific database.

2.6. XAS analyses

XAS analyses of the S K-edge were undertaken at the Diamond Light Source, Oxfordshire, UK on the microfocus spectroscopy beamline I18 with the aim of examining the structural relationship between the sulfur component of the glutathione capping agent and the CdSe nanocrystals. Measurements were taken with an unfocused beam in fluorescence mode with a 4 element Si drifts detector. A range of standards were measured, including oxidized and reduced forms of glutathione, elemental sulfur, sodium sulfite and sodium sulfate. To remove any MOPS buffer, AQDS and unreacted glutathione, GSH–CdSe was concentrated and purified by re-suspension in the nonsolvent 2-propanol as previously described [7, 18].

Further XAS experiments at the Se K-edge (about 12 654 eV) were carried out on the high brilliance x-ray spectroscopy beamline ID26 at the European Synchrotron Radiation Facility (ESRF), Grenoble, France, using a (111) double Si crystal monochromator as described previously [35]. The aim was to examine the differences between the formation of chalcogenide QDs from abiotic and biogenic selenide precursors. In this case, the development of ZnSe with the use of \ensuremath{\beta}-mercaptoethanol (\ensuremath{\beta}ME) as a capping agent (\ensuremath{\beta}ME-ZnSe) was used as an analogue system to CdSe. Owing to the speed of the reaction, which exceeds that which can be seen by QUEXAFS, a novel, in situ, time-resolved experiment was carried out using an injection cell system. The \ensuremath{\beta}ME-ZnSe was synthesized by injecting equal volumes of an anaerobic solution containing 10 mM ZnCl\textsubscript{2} and 24 mM \ensuremath{\beta}ME solution and an anaerobic solution of the appropriate Se^{II-} (5 mM) to produce final concentrations of Zn, Se and \ensuremath{\beta}ME of 5, 2.5 and 12 mM respectively. Initial XAS scans were taken across the Se K-edge after the injection of equal volumes of degassed, deionized water and the abiotic or biogenic Se^{II-} solutions. From these, it was possible to discern two regions of significant difference between the Se^{II-} and ZnSe phases that allowed the reaction and development of ZnSe products to be monitored. The sample cell was then flushed with anaerobic, deionized H\textsubscript{2}O before re-use. The multi element Ge detector was set to analyse at the specified energy with the minimum refresh rate of 0.5 ms during the injection of Zn^{II+}-\ensuremath{\beta}ME and Se^{II-}.

The sample cell was then flushed and repeated for both stated energies for the abiotic and biogenic Se^{II-} solutions.
3. Results and discussion

3.1. Particle characterization and development

The production of glutathione-stabilized cadmium selenide quantum dots by abiotic and biogenic routes was compared. In both cases, the GSH–CdSe suspensions changed colour from yellow to red during refluxing, corresponding to the growth of quantum dots. The fluorescence under UV illumination at a wavelength of 365 nm is shown in figure 1. The UV/vis absorption spectra of these GSH–CdSe aliquots are shown in figure 2. Both abiotic- and biogenic-derived sample sets display primary and secondary absorption peaks for the 0 and 4 min reflux time points; at 356 and 408 nm for the abiotic samples, and at 314 and 334 nm for the biogenic samples. The relative intensity of the secondary, lower wavelength absorption peaks decreased with increasing reflux time for both abiotic and biogenic samples. The peak absorption wavelength for both sets of samples increased with increasing reflux time (figure 2), and a second absorption peak was observed in the abiotic sample sets approximately 30 nm above the primary absorption peak.

To quantify the photoluminescence observed in figure 1, the GSH–CdSe quantum dots were analysed using a constant excitation wavelength of 365 nm and the results are shown in figure 2. The peak emission wavelength for the abiotic GSH–CdSe prior to reflux was at 510 nm, which increased to 590 nm following 20 min of refluxing. The biogenic QDs prior to reflux also had a peak emission wavelength of 510 nm, which increased to 531 nm following refluxing for the same period, 59 nm lower than that of the abiotic QDs. A large shift in emission spectra was observed for the abiotic QDs between 4 and 12 min refluxing time, which was not observed in the biogenic photoluminescence spectra. Secondary emission peaks occurred in the abiotic GSH–CdSe photoluminescence spectra at 117 nm above the peak emission wavelength (figure 2), becoming more prominent with prolonged reflux time. The biogenic spectra initially showed a secondary emission peak occurring 13 nm above the primary emission wavelength (figure 2), but this peak disappeared with subsequent refluxing.
HR-TEM was employed to characterize the particle size, shape and morphology of the QD reflux aliquots (figure 3). Spherical, crystalline CdSe particles with diameters of less than 8 nm were discernable, and lattice fringe measurements corresponded to d-space values of planes within hexagonal CdSe (cadmoselite, ICDD 00-008-0459).

Particle size distributions were calculated using HR-TEM images, and the resulting histograms are shown in figure 4. Abiotic and biogenic QDs initially showed equal average particle diameters of 2.3 nm (±1.3 nm for both biogenic and abiotic). An increase in size distribution was noted, however, for the abiotic QDs after 8 min reflux, with an average particle diameter of 3.2 nm ± 1.2 nm compared to 2.9 nm ± 1.2 nm for the biogenic QDs. Following 20 min reflux, average particle diameters increased to 3.8 nm ± 1.2 nm and 3.6 nm ± 1.2 nm for abiotic and biological QDs, respectively.

Figure 4 shows the peak photoemission wavelength, the first order derivative of the peak photoemission wavelength and the average particle diameter determined by TEM. Using the peak photoemission wavelength as a proxy for particle size, the derivative of this can infer the rate of change of particle growth rate as a factor of temperature at 4 min intervals between sampling. These results show that the rate of change in the peak photoemission wavelength (and accordingly the particle size) increases dramatically above 90°C for the abiotically synthesized QDs, whereas the biogenic QDs display a lower, stable growth rate in this temperature range. PL spectra (figure 2) and particle size distributions from HR-TEM (figure 4) for both abiotic and biogenic QDs show a non-Gaussian size distribution with a tail extending to the high particle sizes.

The results of these analyses show that biogenic Se\(^{II}\) is a viable alternative to abiotically generated Se\(^{II}\) in the
formation of metal selenide quantum dots, alleviating the need for highly toxic and strongly reducing compounds. Analysis of the optical absorption and emission spectra (figure 2) shows that biogenic Se\textsuperscript{II} can be used as a precursor for the synthesis of highly fluorescent nanoparticles comparable to abiotic equivalents. Particle sizing in conjunction with UV/vis and PL emission spectra show that the growth of biogenic QDs occurs at a slower rate than with abiotically generated Se\textsuperscript{II}, suggesting that the presence of bacterially derived moieties, such as proteins, may inhibit the growth rate of CdSe QDs.

The more consistent particle size of the biotic samples could be attributable to a particle size control dictated by coordinating bacterially derived moieties, such as previously reported for sulfurous peptides [11, 15–17, 19, 21]. The development of the large particle size fraction is a deleterious characteristic of aqueous QD synthesis techniques in comparison with organometallic synthesis routes, and although the presence of bacterial proteins leads to an increased control of particle growth by reducing particle growth rate (figure 5), a post-preparative purification step as used by Gaponik et al [7] may be required.

3.2. Protein identification

The impact of secreted bacterial and fungal proteins on the nucleation and growth of metal chalcogenide quantum dots has been observed previously [15, 16, 19, 22, 26, 28]. In order to determine the nature of the bacterially derived moieties in the biogenic Se\textsuperscript{II} solution in this study, both the biogenic and the abiotic Se\textsuperscript{II} solutions were analysed for protein content by SDS-PAGE (figure 6). No protein bands were visible in the abiotic Se\textsuperscript{II} solution; however the red-orange band at the end of the gel for both the biogenic and abiotic samples indicates the precipitation of elemental selenium nanoparticles as a result of oxidation of the selenide solution. The biogenic Se\textsuperscript{II} solution contained a range of proteins of varying mass, of which one was putatively identified as the alpha-subunit of methylmalonyl-CoA decarboxylase, similar to that identified in Veillonella parvula [36], and likely originating from V. atypica.

3.3. XAS investigations

3.3.1. Bonding of the reduced glutathione capping agent.

The coordination environment of sulfur in the reduced glutathione capping agent associated with the CdSe quantum dots was investigated using XAS at the S K-edge; the XANES results are shown in figure 7, along with relevant model compounds. The XANES spectra of the model compounds show a range of spectral shapes and peak energies with an approximately 10 eV difference in peak energy between elemental S and S\textsuperscript{VI}. The spectral profile and absorption edge energy at the S K-edge obtained for the precipitated GSH–CdSe QDs show that S is present in a reduced form, and closely resembles the CdS standard at about 2473 eV (figure 7). There is no evidence for the presence of S–S bonds as in oxidized GSH, or S–O bonds as the oxidized sulfate or sulfite forms. There are two possible Cd–S coordination environments within the samples tested; (i) the Cd–SR bond, where R is the glutathione molecule, between Cd atoms exposed at the surface of the nanoparticles and the S of the cysteine component within glutathione, in which the Cd–S bonds show similar character to those observed for xanthate attachment to ZnSe [37]; and (ii) structural incorporation of free S into a mixed CdSe\textsubscript{1−x}S\textsubscript{x} phase, due to the decomposition of glutathione during reflux in alkaline, aqueous media, as reported for the incorporation of S into CdTe quantum dots [7, 38]. Gaponik et al [7] postulated that excess thiol, coupled with decreasing free Se\textsuperscript{II}– concentrations as particles grow during reflux, increases the proportion of S structurally incorporated, thereby causing an increasing S:Se ratio in the outer layers of the particle. However, the limitation of the QD particle size to below 4 nm is evidence that the Cd–SR bond is restricting particle growth. Differentiating between sulfur attached to...
cadmium at the surface and sulfur structurally incorporated into the outer CdSe layers is not possible using standard XANES techniques. Understanding the relationship between sulfurous capping agents and metal selenide QDs, as well as determining the relative proportions of the Cd–SR and the CdSe$_{1-x}$S$_x$ during particle growth, is challenging and requires collection of an extended x-ray absorption fine structure (EXAFS) time course data set, at the S K-edge, covering QD formation and growth during reflux. This will be the focus of future research.

3.3.2. Selenide stability and ZnSe nucleation. XAS was used to compare the stability of biogenic and abiotic Se$^{II-}$ starting materials, in order to determine the impact of extracellular biological materials (figure 8). As with S, an increase in oxidation state of Se at the K-edge is easily discernable by an increase in absorption edge energy (7.5 eV between Se$^0$ and Se$^{VI}$).

The results show that the initial biogenic Se$^{II-}$ solution was composed predominantly of Se$^{II-}$ with a peak at 12,653.5 eV, but with the possible presence of a minor oxidized component as indicated by the broad bimodal lineshape (figure 8). The stability of the abiotic and biogenic Se$^{II-}$ solutions was compared by repeated scanning across the Se K-edge for 15 min (figure 8). Over this time period, the biogenic Se$^{II-}$ solution remained in a reduced form indicating that it was stable and did not undergo oxidation. However, the abiotic Se$^{II-}$ peak shifted dramatically towards a principally Se$^{VI}$ form, after 15 min exposure to the beam, indicating that the abiotic Se$^{II-}$ was susceptible to oxidation, possibly induced by the x-rays. The proteins identified in the biogenic Se$^{II-}$ solution may coordinate with the Se$^{II-}$ and act as stabilizing agents, limiting oxidation.

The addition of the ZnCl$_2$-βME solution to both biogenic and abiotic Se$^{II-}$ stocks leads to the rapid formation of βME-ZnSe nanocrystals, as shown in figure 9. From these spectra, there is a discernable difference in absorption intensity at 12,661 and 12,667 eV between the Se$^{II-}$ solutions and βME-ZnSe suspensions for both the biogenic and abiotic samples. The change in intensity at these energies defines the rate of nucleation of ZnSe QDs, both in the presence and absence of extracellular biological material (figure 10). First order derivatives of the time-resolved spectra obtained highlight differences in the rate of transformation from the Se$^{II-}$ phase to the ZnSe phase, with the biogenic Se$^{II-}$ samples displaying a wider peak than abiotic counterparts; full-width half-maximum (FWHM) values for these reactions are 1.25 ms and 1.23 ms for the biogenic Se$^{II-}$ samples at 12,661 eV and 12,667 eV, respectively and 0.77 ms and
Figure 9. Se K-edge x-ray absorption spectra for the biogenic (top) and abiotic (bottom) Se\textsuperscript{II} solutions (black lines) and β-mercaptoethanol capped ZnSe (grey lines) formed following addition of ZnCl\textsubscript{2}. The energies used for the time-resolved scans are indicated.

It is possible that proteins present in the biogenic Se\textsuperscript{II}− are also implicated in decreasing the rate of reaction between Zn\textsuperscript{II+} and Se\textsuperscript{II}−, as shown in figure 10 where the FWHM value for two independent biogenic Se\textsuperscript{II}− reactions are twice that for those obtained with abiotic Se\textsuperscript{II}−. However, there are limitations associated with these results, as the FWHM reaction times calculated are of the same magnitude as the minimum detector refresh rate (0.5 ms), although the repetition of the results from two separate biogenic Se\textsuperscript{II}− reactions and two abiotic Se\textsuperscript{II}− reactions suggests these findings are valid; further investigations using ultrafast time-resolved detectors is warranted.

4. Conclusions

The technological, industrial and biological advances using semiconducting, nanoparticulate ‘quantum dots’ merits the investigation of techniques aimed at reducing the economic and environmental costs associated with their manufacture. In this study, we have compared the formation of metal selenide quantum dots from bacterially derived Se\textsuperscript{II}− with quantum dots synthesized from tradition chemical precursors. The use of bacterially derived Se\textsuperscript{II}− in the manufacture of metal selenide quantum dots controls the rate of formation and growth of the QDs, with a decrease in initial reaction rate and a narrower size distribution of particles in comparison to abiotic counterparts, even in the presence of thiol capping agents. The decreased particle size distribution and the 0.74 ms for the abiotic Se\textsuperscript{II}− samples at 12 661 eV and 12 667 eV.

Figure 10. Time-resolved energy scans (main image: first order derivates, inset: time-resolved scans) for (A) biogenic ZnSe at 12 661 eV, (B) biogenic ZnSe at 12 667 eV, (C) abiotic ZnSe at 12 661 eV and (D) abiotic ZnSe at 12 667 eV. Inset: X axis values are time in milliseconds.
reduced particle growth rate allow for tighter size constraints with biotic materials compared to wet synthesis experiments with abiotic Se II−.

Protein gel electrophoresis has shown that bacterial contributions to the synthesis of technologically relevant materials is not limited to direct formation of the Se II− solutions; extracellular biological materials play an important role in Se II− stability after the cells have been removed. S K-edge XANES has supported the hypothesis that GSH acts as a capping agent and is attached to the surface of the QDs. Novel, in situ time-resolved XAS experiments have also demonstrated that the presence of bacterially derived proteins provides increased control over QD size by reducing the rate of initial particle formation.

This research describes an alternative, ‘green’ pathway that utilizes the advantages of biosynthetic techniques, namely the cheap, low temperature and relatively safe synthesis of Se II− without the use of highly toxic and expensive precursors. The results also highlight the stability of the precursors and enhanced control of QD synthesis that can be obtained in the presence of biological coatings. Further research into the characterization and optimization of the role of bacterially derived sulfurous capping agents in metal selenide quantum dot formation is warranted prior to assessing the economic viability and scalability of this technique on an industrial scale.

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