Helmholtz-Zentrum Dresden-Rossendorf (HZDR)



Methyl selenol as precursor in selenite reduction to Se/S species by methane-oxidizing bacteria

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Originally published:

November 2019

Applied and Environmental Microbiology 85(2019)22, e01379-19

DOI: https://doi.org/10.1128/aem.01379-19

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- 1 Selenite Reduction by Methane-oxidizing Bacteria: Methyl Selenol as Precursor for
- 2 Methylated Se/S Species and Characterization of Se_{8-x}S_x Nanoparticles
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- 16 Summary
- 17 A wide range of microorganisms have been shown to transform selenium-containing
- oxyanions to reduced forms of the element, particularly selenium-containing nanoparticles.
- 19 Such reactions are promising for detoxification of environmental contamination and
- 20 production of valuable selenium-containing products such as nanoparticles for application in
- 21 biotechnology. It has previously been shown that aerobic methane-oxidising bacteria,
- 22 including *Methylococcus capsulatus* (Bath), are able to perform methane-driven conversion
- of selenite (SeO₃²) to selenium-containing nanoparticles and methylated selenium species.
- Here, the biotransformation of selenite by Mc. capsulatus (Bath) has been studied in detail

- via a range of imaging, chromatographic and spectroscopic techniques. The results indicate that the nanoparticles are produced extracellularly and have a composition distinct from nanoparticles previously observed from other organisms. The spectroscopic data from the methanotroph-derived nanoparticles are best accounted for by a bulk structure composed primarily of octameric rings in the form Se_{8-x}S_x with an outer coat of cell-derived biomacromolecules. Among a range of volatile methylated selenium and selenium-sulfur species detected, methyl selenol (CH₃SeH) was found only when selenite was the starting material, although selenium nanoparticles (both biogenic and chemically produced) could be transformed into other methylated selenium species. This result is consistent with methyl selenol being an intermediate in methanotroph-mediated biotransformation of selenium to all the methylated and particulate products observed.
- Keywords: Selenite reduction, Elemental selenium, Methane-oxidizing bacteria, Mixed
 chalcogenide amorphous nanoparticles

Introduction

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39 A key biotransformation mechanism of most microorganisms exposed to selenium oxyanions 40 is dissimilatory reduction to nanoparticulate elemental selenium (Lortie et al., 1992; Kessi et 41 al., 1999; Prakash et al., 2009; Nancharaiah & Lens, 2015b). The formation of the 42 nanoparticles (NPs) reduces the toxicity and bioavailability of the selenium species (Combs et 43 al., 1996; Nancharaiah & Lens, 2015a; Eswayah et al., 2016; Song et al., 2017; Vogel et al., 44 2018). Not only does the formation of the NPs reduce the adverse environmental impact of 45 the oxyanions on the microorganisms and their surroundings but present an approach that can 46 potentially be harnessed to produce selenium NPs tailor-made for a variety of technological, 47 clinical, analytical and industrial applications (Prasad, 2009; T. Wang et al., 2010; Bai et al., 48 2011; Tian et al., 2012; Iranifam et al., 2013; Ren et al., 2013; Tran & Webster, 2013; J. 49 Wang et al., 2014; Ramya et al., 2015; Jain et al., 2016; Wadhwani et al., 2016). However, 50 made-to-order NPs with microbial intervention can only be achieved when the structural 51 features of the NPs produced by the different bacteria are better understood and characterised.

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53 In their 2004 paper, Oremland et al investigated the structural features of selenium 54 nanospheres produced by Se-respiring bacteria (Oremland et al., 2004). They found that the 55 three bacteria studied produced red amorphous Se NPs with distinct features, two contained 56 predominantly Se₆ chain units, and the third had Se₈ ring units. Hu and Barton found in an 57 investigation into the reduction of selenium oxyanions by Desulfovibrio desufuricans, a 58 sulfate-reducing bacterium, that amorphous spherical submicro particles containing selenium 59 and sulfur were produced both inside and outside the cell. They found that the bacterium was 60 more effective at reducing selenite. The authors also proposed the mechanisms by which the 61 particles were formed (Hu and Barton, 2013). More recently, Vogel et al proposed that the 62 biogenic selenium NPs produced by Azospirillum brasilense from the biotransformation of 63 selenite is cyclic Se_{8-n}S_n with Se₆S₂ the most likely structure (Vogel et al., 2018). Tugarova et 64 al reported the characterisation of amorphous selenium particles produced by Azospirillum 65 thiophilum and found the presence of only selenium with no evidence of either Se-S or S-S 66 bands in the Raman spectrum (Tugarova et al., 2017). Moreover, Ruiz-Fresneda et al (2018), 67 described the ability of Stenotrophomonas bentonitica for biogenic reduction of Se(IV), 68 production of amorphous Se⁰ (a-Se) nanospheres and their subsequent transformation to one-69 dimensional (1D) trigonal selenium (t-Se) nanostructure where sulfur was associated with the 70 SeNPs. In our study, Eswayah et al, we presented indirect evidence from transmission electron 71 microscopy (TEM) imaging with energy dispersive X-ray spectrometry (EDX) measurements 72 to show that sulfur was associated with amorphous selenium in the extracellular NPs that are 73 produced from the reduction of selenite by methane-oxidizing bacteria (Eswayah et al., 2017). 74

Besides the above-mentioned studies on the biotransformation of selenium oxyanions most investigations have focused on the formation of the NPs. A few others have identified concomitant release of volatile selenium species into the headspace gas (Chasteen, 1993; Burra *et al.*, 2010). However, none of these approaches has provided enough information to enable the elucidation of the processes leading to the formation of the amorphous NPs. Indeed, the formation of the extracellular amorphous NPs forms as reported in many studies (Oremland *et al.*, 2004; T. Wang *et al.*, 2010; Dhanjal & Cameotra, 2010; Zhang *et al.*, 2012; Kamnev *et al.*, 2017) may indicate limited direct involvement of the microorganisms in their formation. In our experiments, we observed the size of the extracellular nanoparticles increases with time, which suggests that the growth of the NPs is a result of abiotic reactions in the culture medium outside the cells. The formation of mixed chalcogenide species by

exchange reactions, when both Se and S species are present in the gaseous and solution phase,

has been reported (Meija & Caruso, 2004; Vriens *et al.*, 2015). It is probable that similar reactions occur in the culture medium solution resulting eventually to the formation of the nanoparticles.

In order to gain a better understanding of the biotic and abiotic transformations occurring in the culture, both selenium- and sulfur-containing species were sampled from the headspace and solution of selenite amended and control samples at fixed times by sorptive extraction in conjunction with analysis by thermal desorption - gas chromatography- mass spectrometry (TD-GC-MS) to identify the compounds. In parallel, the formed NPs were characterised by a range of physical techniques, namely; attenuated total reflectance Fourier transformation infrared spectroscopy (ATR-FTIR), Raman spectroscopy, transmission electron microscopy (TEM) and energy dispersive X-ray (EDX) spectrometry, X-ray absorption spectroscopy (XAS), and X-ray photoelectron spectroscopy (XPS). Herein, the results obtained from these measurements are used to inform the formation and elucidation of the structure of the sulfur-doped red amorphous selenium NPs produced when the methane-oxidizing bacterium *Methylococcus capsulatus* (Bath) reduces selenite.

Results

Preliminary investigations with the methanotroph species Mc.capsulatus (Bath) showed that the sizes of the NPs grew rapidly from an average of 220 ± 51 nm in the first 4 hrs to about 400 ±77 nm in the next 44 hrs as previously published (Eswayah et~al., 2017). It is evident from the HAADF-STEM imaging and TEM thin-section micrographs of the nanoparticles produced by the bacterium that the NPs are associated extracellularly with the cells (see Fig. 1a). Furthermore, Se and S distributions in the EDX maps overlap, indicative of a spatial and likely structural association, suggesting the formation of mixed chalcogenide nanoparticles (see Fig.1b). The intensity of the Se signals was, however, much higher than that for S (see Fig. 1c). In addition, examination of the $S_{k\alpha}$ map reveals that not only was there sulfur in the particles but there was a trail of the element linking the particles to likely sulfur-containing proteins from the bacterial cells.

Se K-edge X-ray Absorption Near-Edge Structure (XANES) spectra of the particles formed by *Mc. capsulatus* (not shown) are in line with those of red amorphous Se with no detectable residual selenite present in the samples. Shell fitting of the Extended X-ray Absorption Fine-

Structure (EXAFS) spectra showed the characteristics of red elemental Se⁰, with a Se-Se path at a distance of 2.35Å. A second Se-Se shell was detected at 3.69Å with coordination numbers less than what has been previously been observed for amorphous Se (Scheinost & Charlet, 2008; Scheinost *et al.*, 2008), possible indication for the presence of a mixed Se/S phase. This small coordination number cannot be attributed to the NP size far above 10 nm, nor to a high structural disorder, the Debye-Waller factor (σ^2) of this second Se-Se showed increasing structural order with particle growth over time, without increasing the respective coordination numbers (see Eswayah et al (2017) for a detailed discussion of these results). The small coordination numbers hence suggest the presence of (more weakly backscattering) S atoms, but attempts to establish their presence by shell fitting failed because of strong destructive interference between Se-Se and Se-S paths.

In order to unravel the make-up of the nanoparticles, the surfaces of the particles were characterized by FTIR, XPS and Raman spectroscopy.

FTIR Analysis

The FTIR spectra of the freeze-dried selenium nanoparticles produced by the *Mc. capsulatus* in liquid NMS medium amended with selenite, the Chem-NPs and the bacterial biomass are shown in Figure 2. The assigned bands are summarised in Table 1.

XPS analysis of the particle surface

The surface composition of the harvested red particles was obtained by XPS, and the elemental content are summarised in Figure 3a. In addition to the selenium which is present at a concentration of 1.25 Atomic weight%, there are five other elements: carbon (46.32%), oxygen (31.41%), nitrogen (8.61%), calcium (5.89%) and phosphorus (4.77%) that were detected on the surface of the particles. The presence of the first three elements is an indication that there are organic molecules on the particle surfaces. The high resolution spectra scans for Se, C, N, and O, and the assigned chemical species from the core level XPS spectra of C1s, N 1s, O 1s bands are shown in Figures 3b-e. The spectrum for 3d Se shows a doublet which is unresolved. However, the fitted deconvoluted peaks show two predominant bands at 55.16 and 56.02 binding energy (eV), and two minor peaks at 55.75 and 56.61 eV, respectively, the later pair of peaks resulting from the Se 3d peak split by spin orbit coupling into Se 3d5/2 and Se 3d3/2. The observed range of binding energies between 55.16 and

- 147 56.61eV is indicative of the presence of reduced selenium species, including elemental
- 148 selenium (Naveau *et al.*, 2007).

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Raman characterisation of the amorphous particles

- Vibrational spectroscopy particularly Raman spectroscopy has been the technique of choice
- 151 for the characterisation of Se_n allotropes, and aggregates. The deconvoluted spectrum
- obtained between 50-600 cm⁻¹ Raman shift from the harvested SeNPs is shown in Fig 4.
- There are four bands which are visible: the main band at 251.5 cm⁻¹ and smaller ones at 80.2,
- 358.8 and 506.5 cm⁻¹, respectively. All of the bands were present in all of the scans of
- samples collected at different time points: 6, 24 and 48h. The band at 513.5 cm⁻¹ was more
- prominent than that at 358.8 cm⁻¹. The band at 80.2 cm⁻¹, which is a shoulder, is only visible
- in the deconvoluted spectrum.

Speciation of selenium and sulfur in the medium solution and headspace

- The GC-MS chromatograms of the species found in both the headspace and solution of the
- selenite amended medium at 4h and 20h are shown in Figures 5 and 6, respectively. The
- earlier time was chosen because the formation of the particles and therefore the red colour of
- the solution were barely discernible. A summary of the selenium- and sulfur-containing
- species are given in Table 3. Examination of the data in the Tables showed that after 4h, three
- 164 compounds: methyl selenol (MSeH), dimethyl selenenyl sulfide (DMSeS) and dimethyl
- diselenide (DMDSe) were detected in both the solution and headspace. In addition dimethyl
- diselenenyl sulfide (DMDSeS) was also found in the solution. At 20h, three new species:
- Bis(methylselenomethane) (DMSe), dimethyl selenenyl disulfide (DMSeDS), and dimethyl
- diselenenyl sulfide (DMDSeS) were detected in the solution in addition to triselenothone/
- dimethyltriselenide (DMTSe) which was detected in both the solution and headspace.

Discussion

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- 171 The results from the kinetics experiments showed that there were increases in particles sizes
- with incubation time leading us to hypothesize that much of the structure of the particles was
- formed in the extracellular space. If this is true, then the key reactions resulting in the
- increase in the particle sizes are essentially abiotic in nature. Consequently, the clues to the
- structural formation of the particles must lie in the nature and identity of the compounds that
- are concomitant in the solution and headspace of the nascent particles. It was therefore

essential to sample for selenium- and sulfur-containing compounds in both the headspace and solution, followed by their analyses and identification.

The FTIR spectra of the SeNPs produced by *Mc. capsulatus*, samples of biomass of the strain (control), as well as the Chem-SeNPs were recorded in order to identify the functional groups capping the synthesized SeNPs. The peak centred at 3297 cm⁻¹ corresponds to the –OH and –NH stretching vibrations of the amine and carboxylic groups. Peaks at 2927 cm⁻¹ corresponded to the aliphatic saturated C–H stretching modes (Naumann *et al.* 1995; Kamnev *et al.* 2017). The peaks at 1644, 1538, and 1239 cm⁻¹ are characteristic of amide I, amide II, and amide III bands of proteins, respectively (Alvarez-Ordóñez *et al.* 2011; Ojeda & Dittrich 2012). The symmetrical stretch of carboxylate group can be attributed to the bands observed at 1366 cm⁻¹. The peaks at 1150, 1077 and 1015 cm⁻¹ corresponded to the C–O stretching vibrations of C-O-C groups (Naumann *et al.* 1995; Beekes *et al.* 2007). The presence of phosphoryl groups was confirmed by the peak at 919 cm⁻¹. Additionally, peaks at 859 and 762 cm⁻¹ (fingerprint region) could be mainly attributed to aromatic ring vibrations of aromatic amino acids (tyrosine, tryptophan, phenylalanine) and possibly nucleotides (Burattini *et al.* 2008; Kamnev 2008).

The FTIR spectra of SeNPs of *Mc. capsulatus* differ from those of the bacterial biomass (control) and the Chem-SeNPs. The main difference between the spectra is that the Bio-SeNPs exhibit more peaks in the protein and polysaccharide vibration region, indicating the presence of proteins and polysaccharides in the biomacromolecules capping the SeNPs (Shirsat *et al.*, 2015; Wadhwani *et al.*, 2016; Tugarova & Kamnev, 2017; Kamnev *et al.*, 2017).

By contrast, Chem-SeNPs obtained through reaction of Na₂SeO₃ with L-cysteine displayed a broad absorption band around 3350 cm⁻¹ and absorption band at 2923 cm⁻¹ that are assigned to O–H vibrations of the absorbed H₂O and C–H vibration in the alkyl chain of L-cys, respectively. The peak at 1606 cm⁻¹can be mainly attributed to C=O vibrations. It is noteworthy that the presence of organic residues such as carbohydrates, lipids, and proteins on the surface of biogenic SeNPs were completely absent in the Chem-SeNPs spectrum (see Figure 2) FTIR spectra of the Bio-SeNPs separated from the *Mc. capsulatus* cells showed bands typical of proteins, polysaccharides and lipids associated with the particles (in line with their TEM images showing a thin layer over the particles), in addition to strong carboxylate bands, which may stabilise the SeNPs structure and morphology.

The XPS results show two Se containing species, and other organic constituents.

In assigning the Se bands, it is essential to link the structure of the particles to the selenium-containing species that have been identified in the solution (see the discussion of the TD-GC-MS results below). Since data for the exact compounds are not available, structures that may be similar to these in the particles have been selected. The major band at 55.16 eV, has been assigned to the compound: (CH₃)₂NC(Se)SeC(Se)N(CH₃)₂ (Kobayashi *et al.*, 1986). This is a reasonable fit to the results obtained in this study not only because of the presence of selenium but also the content of the methyl groups and nitrogen. The nitrogen could account for the single band at 400.10eV usually assigned to amine nitrogen. The bands at 56.02 eV has been assigned to (-CSeC(CH₃)C(CH₃)Se-)2 (Dáaz *et al.*, 1996). Both of these assignments are an indication of the presence of long chain of selenium-containing methylated species. Missing from both spectra is oxygen which is presumably present as C=O, and either C-O-C or C-O-H at 531.58 and 532.84 eV, respectively.

However, the presence of Se-C was not detected in the spectrum thus indicating the amount of carbon directly bound to selenium was low. Furthermore, because of the high concentration of selenium in relation to sulfur, the signal for the latter could not be resolved and identified.

Whereas, both FTIR and XPS provided information on the forms in which the elements are present on the particle surface, Raman spectroscopy enables the identification of the basic structural make-up of the particles. It has been proposed that amorphous selenium is composed of a mixture of Se_n rings and helicoidal chains (Lucovsky *et al.*, 1967; Yannopoulos & Andrikopoulos, 2004; Demchenko *et al.*, 2010). The proportion of each depends on chemical and physical conditions under which the samples are made, and the treatments to which they have been subjected. According to Carini *et al* the band at around 250 cm⁻¹ is characteristic of amorphous selenium (Carini *et al.*, 1980). The symmetrical band at 251.5 cm⁻¹, with full width half maxima of 30 cm⁻¹ found in this study is characteristic of Se-Se stretching vibration in pure Se₈ (Lucovsky *et al.*, 1967; Baganich *et al.*, 1991; Nagels *et al.*, 1995) with its' deformation vibration at the deconvoluted band of 80.2 cm⁻¹. However, in the study of mixed selenium and sulfur alloys by Machado et al, the authors state that as the sulfur concentration increases in mixed amorphous selenium, the peak at 234 cm⁻¹ usually associated with Se chains decreases in intensity and the band at 250 cm⁻¹, usually associated with selenium rings increases in intensity. They also observed the appearance of a band at

- 352 cm⁻¹ as the selenium to sulfur ratio increases to either 4:1 or 7:3 (Machado *et al.*, 2010).
- The band at 358.8 cm⁻¹ has been assigned to S-Se stretching vibration (Eysel & Sunder, 1979;
- 243 Kasuya et al., 1996). The band at 513.5 cm⁻¹ is probably due to the Se-Se overtone band of
- 244 the fundamental band at 251.5cm⁻¹. An important observation in this regard is that the S-S
- vibration band is not seen when the S composition is below that found in Se₆S₂ (Lucovsky *et*
- al., 1967). Therefore it is probable that the composition of the harvested particles is Se_{8-x} , S_x ,
- were x is equal to or greater than 2.
- The intensity of the band 251.5 cm⁻¹ is indicative of the predominance of the Se-Se bonds in
- the structure of the particles. It is probable that S is integral to the mixed particle structure
- Se_{8-x}S $_{x}$, and not as an S₈ impurity in the particles. The low available sulfur content in the
- medium makes the formation of an S-S homonuclear bond in the mixed particle structure
- 252 highly unlikely (Eysel & Sunder, 1979).
- We have previously reported that methyl selenol is produced and detected in the head space
- only when selenite is in the starting medium (Eswayah et al, 2017). Besides the presence of
- methyl selenol in the headspace, methyl selenoacetate was also detected. However, in the
- present study methyl selenoacetate was not detected when the headspace sorptive extraction
- probes were deployed for sample collection. No methyl selenol was detected when either the
- harvested or chemically synthesized nanoparticles were added to the medium in the absence
- of selenite. Therefore, we propose that methyl selenol may be the precursor of all of the
- methylated selenium species as well as the selenium-containing nanoparticles. If this is the
- 261
- case, the first step in the biotransformation of selenite would involve the reduction and
- methylation of selenite to methyl selenol followed by the formation of the other selenium-
- and sulfur-containing species. Indeed, the formation of some of the latter species requires the
- presence of the nascent selenium particles.
- Based on these observations a possible pathway for the formation of the particles and the
- other products of biotransformation of selenite can be outline as in Scheme 1.
- A series of abiotic reactions proposed by Ganther (1968; 1971) and outlined by Xu and Barton
- (2013) implicate glutathione (GSH) and GSH reductase in the production of elemental
- selenium. The proposed steps leading up to the formation of methyl selenol are shown in
- Scheme 2.

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The presence of sulfur-containing species was detected in the headspace and culture supernatant of the control *Mc. capsulatus* culture without added selenite. These species included: benzothiazole, dodecanethiol, and propanesulfonyl, which are the likely source of sulfur in the structure of the particles. Evidence of the formation of the longer chains of the selenium- containing species can be seen in the nature of the compounds that are found in the solution after hours of incubation, first after 4h, dimethyl diselenenyl sulfide was detected, and subsequently dimethylselenodisulfide, dimethyltriselenide and bis(methylseleno) methane were detected after 20h. More complex mixed Se and S compounds are formed in the medium with time. The formation of the mixed chalcogenides of Se and S is hardly surprising since S may be available from the reduction of sulfate in the growth medium. A key question therefore is how the solution chemistry relates to the observed structural features of the Se particulates.

It is likely that these longer chains polymerize to form Se_x or Se_{8-x}S_x linear or cyclic structures. Indeed, all chalogen elements have the tendency to form cyclic allotropes. The dominant allotrope will depend on the experimental conditions. Examples of exchange reactions that may occur based on the presence of the detected selenium- and sulfur-containing species are shown in the following equations:

290 2-Se-Se-S- + -Se-Se- ---->
$$Se_6S_2$$
 (3)

As can be seen a variety of nanocomposites can be formed. Similar exchange reactions have been shown to occur when mixed Se and S complexes are present in the same solution (Vriens et al., 2015). Indeed these reactions are known to occur in amorphous selenium semiconductors (Steudel, 1986). The Raman results indicate these are not open chains clusters but cyclic structures with the Se₈ structure dominant and the probable presence of small amounts of Se-S bonds. To prevent the introduction of sulfur into the structures, the presence of the element could be reduced from amount in the culture medium if this does not affect the bacterial growth.

The formed particles are surrounded and stabilized by the presence of polymeric substances as proposed by Jain *et al* (2015) as evidenced by the XPS and FTIR measurements.

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In this study, we demonstrate for the first time that in the reduction of selenite by Mc. capsulatus (Bath), a methane oxidizing bacterium, methyl selenol is the likely precursor for the formation of methylated selenium-containing and mixed chalcogenides species. Subsequent exchange reactions between the species result in the formation of the amorphous allotropic form of selenium, cyclic Se_8 with sulfur in its structure. The nature of the molecular mediators in reduction of selenite, supplying sulfur that is integral to the structure of the nanoparticles and supplying the methyl groups found in the volatile selenium containing products remain to be identified.

Experimental Procedures

Bacterial strains and growth conditions

The methanotrophic bacterium *Methylococcus capsulatus* (Bath) (NCIBM 11132) was grown and propagated aerobically using methane as the carbon and energy source as previously described (Eswayah et al, 2017). For these experiments the initial selenite concentration used was 20 mg L^{-1} .

Detection of solution and volatile selenium species

Solution and volatile selenium-containing species were sampled by immersive sorptive extraction using sampling probes (HiSorb probe, Markes International, UK) from either the solution or headspace. Extension screw-on arms were fabricated for each probe so that they could be inserted through the Suba-Seals used to seal the necks of the culture flasks. To ensure that the probes and tubes were contamination free, before use, the probes and tubes were preconditioned with helium at flow rate of 90 mL min⁻¹ using the following temperature programme: 15 min at 100 °C, 15 min at 200 °C, 15 min at 300 °C and 15 min at 335 °C. The preconditioned probes were inserted into either the liquid or headspace of the Mc. capsulatus (Bath) culture medium through the Suba-Seals. The probes were removed from the Suba-Seals after different incubation time (4 and 20 h, respectively), rinsed with HPLC grade water, dried with lint-free tissue, and then placed into the thermal desorption tubes (Markes International, UK). Samples analyses were performed on a combined thermal desorption GC–MS system. The volatiles were desorbed at 250°C and concentrated on a thermal desorber (Unity®, Markes International Limited) at -10°C cold trap for 5 min (helium flow 50 mL min⁻¹) and then were transferred onto the GC/MS system (7890A-GC with 5975C-MS, Agilent Technologies) equipped with a capillary column (Agilent J&W HP- MS GC Column, 30 m, 0.25 mm, 0.25

333 μm). Helium was used as the carrier gas at a flow rate of 1 mL min⁻¹, injector temperature,

334 250°C, and the chromatogram was obtained using the following temperature programme: 35°

C for 1 min; 10°C min⁻¹ to 25 0°C; and then held at 25 0°C for 1 min. The National Institute of

Standards and Technology (NIST) MS search program (version 2011) was used to identify the

compounds based on their MS spectrum.

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Extraction of selenium nanoparticles produced by Mc. capsulatus (Bath)

Freshly grown cultures (at OD_{600} of 0.5-0.8) were supplemented with 20 mg L^{-1} SeO_3^{-2} and incubation was continued at 45°C with shaking in the presence of methane. After 48h the development of the reddish colour had occurred, the cultures were pelleted by centrifugation (at $12,500 \times g$; 10 min). SeNPs were extracted by a modification of the method published by Sonkusre et al. (2014), as follows. The resultant pellet was washed and re-suspended in 10 mL of sterile water followed by addition of lysozyme to give a final concentration of 500 μg mL⁻¹, and the tube was incubated at 37°C for 3 h. The suspension was passed through a French pressure cell (1500 psi, 4°C). The resultant slurry containing both cell debris and NPs was washed four times at 15,000 × g for 10 min with 1.5 M Tris-HCl (pH 8.3) containing 1% sodium dodecyl sulfate (SDS). The resultant pellet containing SeNPs and the insoluble cell wall fraction was washed and resuspended in 4 mL sterile water in a 15 mL Falcon tube, and 2 mL of 1-octanol were added. The solution was mixed vigorously on a vortex mixture for five min and centrifuged at 2000 × g for 5 min at 4°C. The tubes were then kept undisturbed at 4° C for 24 hours. The upper phase and interface containing the insoluble cell fraction were removed, and the bottom water phase containing SeNPs was transferred to a clean 15 mL centrifuge tube. This was washed sequentially with chloroform, absolute ethanol, 70% ethanol, and water at 16000 \times g. Collected NPs were re-suspended in water and stored at 4°C. Transmission electron microscopy (TEM) and energy dispersive X-ray (EDX) spectrometry/high-angle annular dark-field (HAADF) scanning TEM (STEM) analysis Samples of selenite amended culture were treated and analysed as previously described

Oxford Instruments XMax SDD EDX detector and a high-angle annular dark-field (HAADF) scanning TEM (STEM) detector. For thin section analysis, after the ethanol dehydration steps, the cells were embedded in EM

(Eswayah et al, 2017). The samples were examined in an FEI Tecnai F20 field emission gun

(FEG)-TEM operating at 200 kV and fitted with a Gatan Orius SC600A CCD camera, an

bed 812 epoxy resin and cut into thin sections (90 nm, using a diamond knife on a Reichert

366	carbon. TEM specimen holders were cleaned by plasma prior to TEM analysis to minimize
367	contamination. Samples were examined with a high-resolution Philips CM 200 transmission
368	electron microscope at an acceleration voltage of 200 kV under standard operating conditions
369	with the liquid nitrogen anti-contaminator in place.
370	X-ray absorption spectroscopy
371	The conditions for the X-ray absorption spectroscopy measurements were as described
372	previously (Eswayah et al, 2017).
373	X-ray photoelectron spectroscopy (XPS) analysis
374	Harvested SeNPs samples were deposited on silicon wafer, left to dehydrate in the load lock of
375	the XPS instrument overnight. The analyses were carried out using a Kratos Axis Ultra DLD
376	instrument with the monochromated aluminium source. Survey scans were collected
377	between 1200 to 0eV binding energy, at 160 eV pass energy and 1 eV intervals. High-
378	resolution C 1s, N 1s, O 1s, Se 3d and S 2p spectra were collected over an appropriate energy
379	range at 20 eV pass energy and 0.1 eV intervals. The analysis area was 700 μm by 300 μm.
380	Two areas were analysed for each sample, collecting the data in duplicate. Charge
381	neutralisation was used with intention of preventing excessive charging of the samples during
382	analysis. The data collected were calibrated in intensity using a transmission function
383	characteristic of the instrument (determined using software from NPL) to make the values
384	instrument independent. The data can then be quantified using theoretical Schofield relative
385	sensitivity factors. The data were calibrated for binding energy using the main carbon peak C
386	1s at 285.0 as the reference peak, and correcting all data for each sample analysis accordingly.
387	Raman spectroscopy analysis of SeNPs
388	Aliquots of 2 µL of SeNPs suspended in water were transferred onto a calcium fluoride (CaF ₂)
389	slide and air-dried prior to Raman analysis. Raman spectra were obtained using a Horiba
390	LabRam HR and a modified Horiba LabRam HR (Wellsens Biotech. Ltd., China). Three
391	factors have been modified in this new Raman system to improve Raman spectral quality.
392	These comprise shortening the Raman light path, employing a low noise and sensitive EMCCD
393	for the Raman signal detection, and increasing incident laser power. The old and new modified
394	systems are identical except these three factors. The Raman signals were collected by a
395	Newton EMCCD (DU970N-BV, Andor, UK) utilizing a 1600 × 200 array of 16 μm pixels
396	with thermoelectric cooling down to -70°C for negligible dark current. A 532 nm Nd:YAG
397	laser (Ventus, Laser Quantum Ltd., UK) was used as the light source for Raman measurement.
398	A 100× magnifying dry objective (NA = 0.90, Olympus, UK) was used for sample observation
399	and Raman signal acquisition. A 600 line/mm grating was used for

400	the measurements, resulting in a spectral resolution of $\sim 1 \text{ cm}^{-1}$ with 1581 data points. The
401	laser power on sample was measured by a laser power meter (Coherent Ltd.). The Raman
402	spectra were processed by background subtraction (using spectra from cell free region on the
403	same slide) and normalization using the Labspec5 software (HORIBA Jobin Yvon Ltd., UK).
404	Fourier transformation infrared (FT-IR) spectroscopy measurements of SeNPs
405	In order to determine the functional groups present on the SeNPs, the FTIR spectra of SeNPs
406	were recorded on a PerkinElmer Spectrum 100 FT-IR Spectrometer equipped with an
407	attenuated total reflectance (ATR) attachment. Spectra were recorded from 4,000 to 650 cm ⁻¹ ,
408	and 4 scans were averaged at a resolution of 4 cm ⁻¹ . Extracted SeNPs were freeze dried
409	overnight and analyzed without further treatment. For comparison, the FTIR spectra of
410	samples of bacterial cells (as control) and chemically synthesized SeNPs (Chem-SeNPs) were
411	also recorded. For the controls, freshly grown cultures ($OD_{600} \sim 0.7$) of <i>Mc. capsulatus</i> (Bath)
412	were centrifuged at $11000 \times g$ for 10 min to obtain the cell pellets. The pellets were washed
413	twice with phosphate buffered saline (Sodium chloride, 150 mM, and sodium phosphate, 150
414	mM) pH 7.2, and then freeze dried overnight. The synthesis of Chem-SeNPs was done
415	according to the procedure of (Lampis <i>et al.</i> , 2017) as follows: 1.0 mL of 50 mM L-cysteine
416	(Sigma-Aldrich, Dorset, UK) solution was added dropwise into 1.0 mL of 0.1 M Na ₂ SeO ₃ .
417	The mixed solution was then stirred for 30 min at room temperature. The Chem-SeNPs were
418	pelleted by centrifugation (at $15000 \times g$; 10 min), and then freeze dried overnight.
419	peneted by centinugation (at 13000 × g, 10 mm), and then neeze direct overlight.
420	
420	Acknowledgement
421	ASE is grateful for the award of a PhD scholarship from the Government of Libya.
422	Conflict of Interest
423	The authors declare no conflict of interest.
424	Originality-Significance Statement
425 426	We certify that all of the research and the conclusions are original and have not been presented elsewhere.
427	

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List of Figures

Figure1 TEM thin-section micrographs of *Mc. capsulatus* (a) exposed to 20 mg L⁻¹ SeO₃²⁻, showing the extracellular locations of the Se⁰ nanospheres, HAADF-STEM imaging, showing Se nanospheres associated with the cells with EDX maps (generated from spectra collected from the indicated areas) of Se and S(b) and TEM of *Mc. capsulatus* cultures exposed to 20 mg L⁻¹ SeO₃²⁻ (c) with EDX analysis within the electron dense regions (Se⁰ nanospheres). Cells were fixed with 3% glutaraldehyde and 2% OsO₄ immediately before the analysis.

Figure 2 The FTIR spectra of freeze dried Bio-SeNPs (blue) and bacterial biomass (red) of *Mc. capsulatus* exposed to 20 mg L⁻¹ SeO₃²⁻ and harvested at $OD_{600} \sim 0.7$, separated by centrifugation, washed with phosphate buffered saline pH 7.2 and freeze dried; as well as Chem-SeNPs (black) obtained through reaction of Na₂SeO₃ with L-cysteine. The spectra are representatives of 5 runs of the experiments.

Figure 3 Wide scan X-ray photoelectron spectra of the SeNPs produced by *Mc. capsulatus* (a) and high resolution spectra for Se 3d, C 1s, O 1s and N 1s are shown in b, c, d and e, respectively. The spectra are representatives of 2 runs of the experiments.

Figure 4 The Raman spectra of purified Se nanospheres from Mc. capsulatus

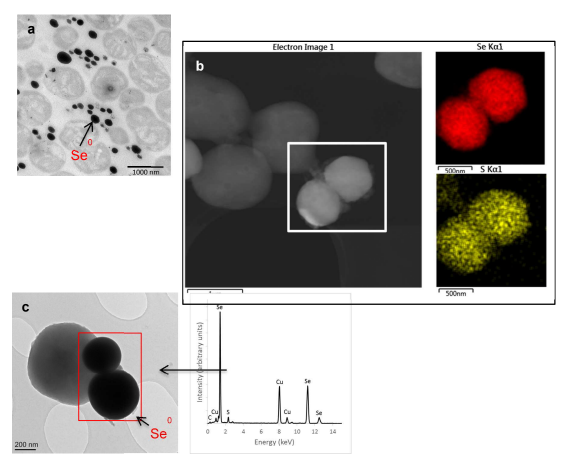
Figure 5 GC mass chromatograms of the liquid phase of the *Mc. capsulatus* (Bath) cultures amended with selenite (20 mg L⁻¹) at 4h (a) and 20h (b). The chromatograms were obtained by selecting the 80 m/z ion, and peak identification was achieved using the GC-MS library.

Figure 6 GC mass chromatograms of the headspace of the *Mc. capsulatus* (Bath) cultures amended with selenite (20 mg L⁻¹) at 4h (a) and 20h (b). The chromatograms were obtained by selecting the 80 m/z ion, and peak identification was achieved using the GC-MS library.

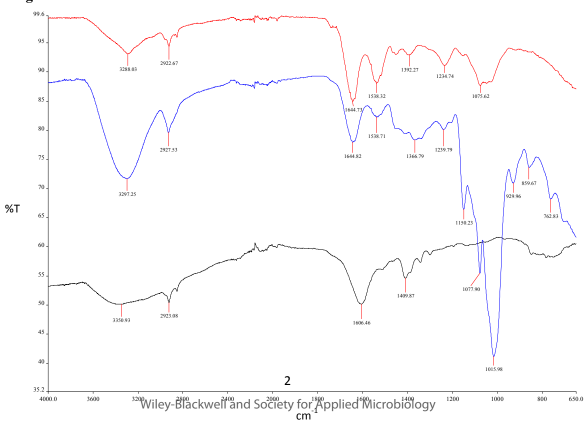
Scheme 1 A schematic diagram showing the reduction of selenite to methyl selenol with the subsequent formation of other selenium-containing species. The numbers donate the following: 1. reduction & methylation, 2. reduction & methylation, 3. polymerization, 4. exchange reactions.

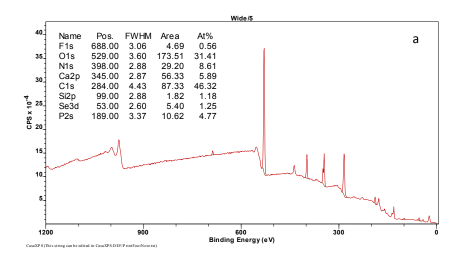
Scheme 2 A schematic diagram showing the formation of methyl selenol, selenium particles and methylated derivatives.

Figure 1









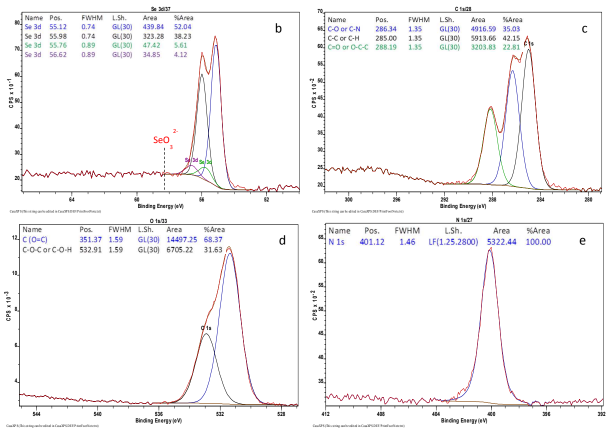
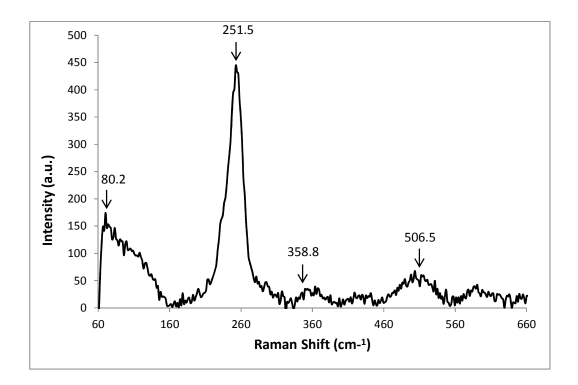
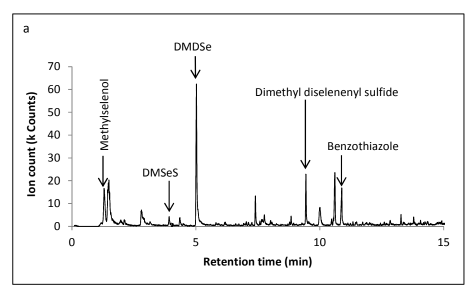


Figure 3

Figure 4





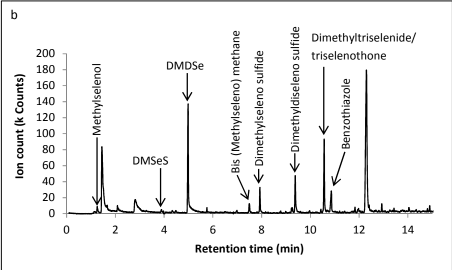
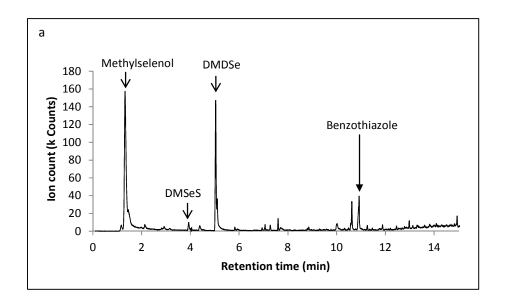


Figure 5



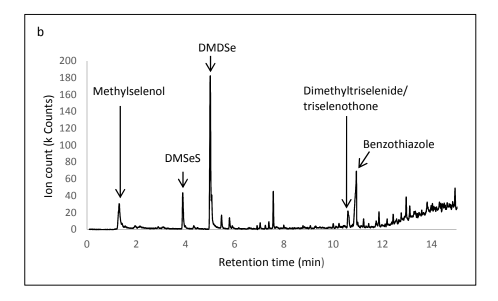
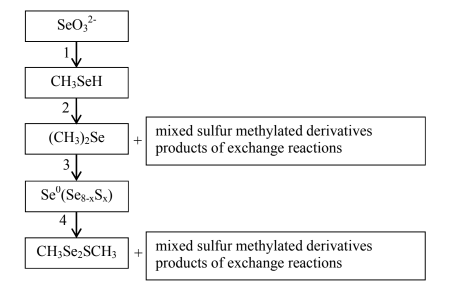
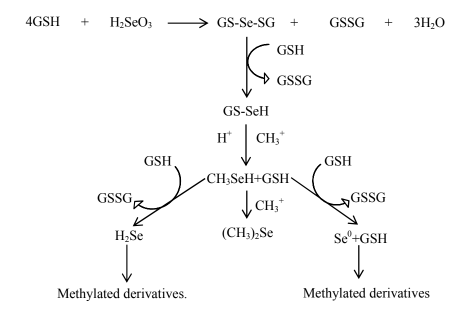


Figure 6

Scheme 1



Scheme 2



List of Tables

Table 1 Tentative assignments of main bands to the relevant functional groups (wavenumber, cm⁻¹) (Naumann *et al.*, 1995; Beekes *et al.*, 2007; Burattini *et al.*, 2008; Kamnev, 2008; Alvarez-Ordonez *et al.*, 2011; Ojeda & Dittrich, 2012; Kamnev *et al.*, 2017).

Sample	O—H; N—H (amide A in proteins)	C—H in >CH ₂)	C=O (ester moiety)	Amide I (in proteins)	Carboxyl COO')	Amide II (in proteins)	-CH ₂ /-CH ₃ (in proteins, lipids, polyesters, etc.)	C=O of COO)	C—O—C/C—C—O (in ester moieties)	Amide III / O-P=O	C-O, C-C, C-H, C-O-C in polysaccharides, and	Phosphoryl groups	"fingerprint region"
Cell biomass of Mc. capsulatus	3288	2922		1644		1538		1392		1234	1075		
SeNPs produced by <i>Mc. capsulatus</i>	3297	2927		1644		1538		1366		1239	1150	919	859
											1077		762
											1015		
Chem-SeNPs		2923			1606			1409					

Table 2. A summary of the selenium- and sulfur-containing species detected in the headspace and solution after 4h and 20h incubation of *Mc. capsulatus* (Bath) in selenite amended medium using sorptive extraction in conjunction TD-GC-MS analysis

Incubation Time		Species										
		Methyl selenol	DMSeS	DMDSe	Bis (Methylseleno) methane	Dimethyl selenosulfide	Dimethyl diselenenyl sulfide	Triselenothone/ Dimethyltriselenide	Benzothiazole	Diethyl sulfuxide	Propanesulfonyl	Dodecanethiol
In solution	4 h	+	+	+	+	-	+	-	+	-	-	-
(selenite amended)	20 h	+	+	+	+	+	+	+	+	+	-	-
In headspace	4 h	+	+	+	-	-	-	-	+	-	-	-
(selenite amended)	20 h	+	+	+	-	-	-	+	+	-	-	-
(Control)	20 h	-	-	-	-	-	-	-	+	-	+	+

^{+ =} detected; - = unknown