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Article

A high resolution study of dynamic changes of Ce2O3 and CeO2 nanoparticles in complex environmental media

Ruth Corrin Merrifield, Kenton P. Arkill, Richard E. Palmer, and Jamie R Lead

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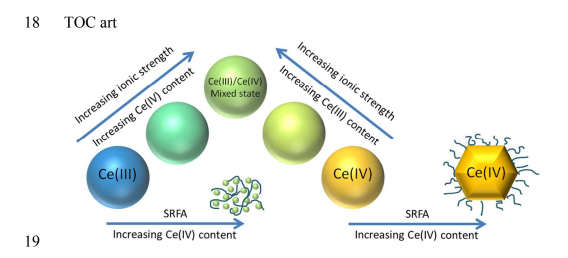
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| 1 | A high resolution study of dynamic changes of Ce_2O_3 and |
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| 2 | CeO ₂ nanoparticles in complex environmental media |
| 3 | |
| 4 5 | Ruth C. Merrifield, ^{1, 2} , Kenton P. Arkill ^{3, 4} , Richard E. Palmer ⁵ , [*] Jamie R. Lead ^{1, 2} |
| 6 7 | ¹ Department of Geography, Earth and Environmental sciences, University of Birmingham, Birmingham, UK |
| 8 9 | ² Center for Environmental Nanoscience and Risk, University of South Carolina, Columbia, SC, USA |
| 10 11 | ³ School of Medicine, University of Nottingham, Nottingham, UK ⁴ CSIC UPV/EHU and PiE, University of the Basque Country, Spain |
| 12 13 | ⁵ Nanoscale Physics Research Laboratory, Physics and Astronomy, University of Birmingham, Birmingham, UK |
| 14 15 16 17 | |



20 ABSTRACT

21 Ceria nanoparticles (NPs) rapidly and easily cycle between Ce(III) and Ce(IV) oxidation states, 22 making them prime candidates for commercial and other applications. Increased commercial 23 use has resulted in increased discharge to the environment and increased associated risk. Once 24 in complex media such as environmental waters or toxicology exposure media, the same redox 25 transformations can occur, causing altered behavior and effects compared to the pristine NPs. 26 This study used high resolution scanning transmission electron microscopy and electron energy 27 loss spectroscopy to investigate changes in structure and oxidation state of small, polymer-28 coated ceria suspensions in complex media. NPs initially in either the III or IV oxidation states, 29 but otherwise identical, were used. Ce(IV) NPs were changed to mixed (III, IV) NPs at high 30 ionic strengths, while the presence of natural organic macromolecules (NOM) stabilized the 31 oxidation state and increased crystallinity. The Ce(III) NPs remained as Ce(III) at high ionic 32 strengths, but were modified by the presence of NOM, causing reduced crystallinity and 33 degradation of the NPs. Subtle changes to NP properties upon addition to environmental or 34 ecotoxicology media suggest that there may be small but important effects on fate and effects 35 of NPs compared to their pristine form.

37 INTRODUCTION

Interactions and transformations of NPs with environmental and toxicological media are 38 known to occur¹⁻³ but are often not well studied, although published work focuses on changes 39 in dissolution, aggregation, shape, size and size distribution³. In the case of ceria NPs that do 40 not readily dissolve in water but can transform chemically⁴, transformations such as Ce(III) to 41 42 Ce(IV) cycling and associated changes are likely to be of key importance for understanding biological and environmental behavior.⁵ These same chemical transformations make ceria NPs 43 of particular interest to industry, in particular they are widely used in microelectronics/semi-44 conductor industries, as mechanical polishers, ⁶⁻⁹ and as a fuel additive in diesel.^{10, 11} The 45 oxidation state of ceria NPs is known to be size dependent ^{12, 13} with the larger particles 46 47 preferentially in the tetravalent (Ce(IV)) state and smaller NPs preferentially in the trivalent 48 (Ce(III)) state. However, particles can easily cycle between the trivalent and tetravalent states of Ce.¹⁴ with implications for biological behavior such as the production of reactive oxygen 49 species (ROS) and oxidative stress. A number of ceria NP types have been shown to have an 50 51 anti- or pro-oxidant effect, but these potentially protective properties are strongly dependent on the surrounding buffer composition and pH¹⁵, while polymer coatings can reduce the toxic 52 effects of the NPs¹⁶. The toxicity of ceria NPs reported in the literature is inconclusive with 53 both low ¹⁷ and higher ¹⁸ toxicities reported. Differences are possibly linked to morphological 54 changes or surface oxidation state, which themselves have been linked to high toxicity via the 55 production of ROS¹². Some studies have also shown that redox behavior can impact 56 dissolution; although ceria is believed to have a low solubility, in some cases ionic Ce from 57 dissolution has been shown to be sufficient to explain some toxicity ¹⁹. 58

60

One key aspect in understanding and then predicting the likely environmental toxicity and behaviors of ceria NPs is their redox (and crystallinity) changes and how this is affected by environmental and toxicology exposure media. This study aims to observe the effect of ionic strength and NOM from representative toxicology exposure and environmental media on the structure and oxidation state of two small, PVP capped ceria NPs. The results will increase understanding in processes such as toxicity and environmental transport.

67

69 MATERIALS AND METHODS

Nanoparticle synthesis. Cerium(III) oxide NPs, nominally 5 nm, were synthesized and 70 characterized as previously published ⁴. Briefly, Ce(III)NO₃ (Sigma Aldrich) was dissolved in 71 72 a solution of 10 KDa polyvinylpyrrolidone (PVP, Sigma Aldrich). The mixture was heated for 73 3 hours at 105 °C, followed by quenching the flask into cold water. When the reaction mixture 74 was cooled, the excess PVP was removed by adding acetone, centrifuging at 4000 rpm for 10 75 minutes using an Eppendorf 5810R bench top centrifuge. The yellow pellet was retained, and 76 the excess liquid was discarded. The pellet was resuspended in ultrahigh purity water (UHP, 77 resistivity 18.2 M Ω cm, total organic carbon <10 ppb). This procedure was repeated three 78 times to ensure all excess PVP was removed. After the final resuspension in UHP water, the 79 NPs were re-dispersed to 100 ml of UHP water and filtered through a 100 nm mixed cellulose 80 ester membrane (EMD Millipore[™]) filter using an asepic vacuum filter system (EMD 81 MilliporeTM) to a selected concentration and stored at room temperature in the dark.

82

Cerium (IV) oxide NPs were produced by converting 50 mL of cerium (III) stock NP 83 84 suspension. 50ml of the Ce₂O₃ NP suspension was placed into a 125 mL Teflon-lined 85 autoclave with 1ml of 1M NaOH and heated at 140 °C for 4 hours. The resulting solution 86 turned bright orange and contained CeO_2 NPs. The suspension was ultrafiltered using an 87 AmiconTM stirred ultrafiltration cell (EMD MilliporeTM) and a 3 KDa UltracelTM ultrifiltration 88 disc, made from regenerated cellulose (EDM MilliporeTM), to remove any excess cerium ions 89 and unreacted NaOH. Finally the suspension was filtered through a 100 nm mixed cellulose 90 ester membrane (Millipore) filter using a vacuum filter system (Millipore) to remove any 91 aggregated particles and stored under the same conditions as the Ce(III) oxide NPs.

93 **Preparation of Ceria NPs in media.** The two NP suspensions were separately added to four different environmental conditions including OECD and ISO algae and daphnia media and a 94 synthetic EPA soft water, and synthetic EPA soft water with 8 mg L^{-1} of Suwanee River fulvic 95 96 acid (SRFA). A full list of conditions, sample abbreviations and their compositions are given in 97 S1 (Table S1-A and S1-B). The particle solutions were left for 72 hours, equivalent to toxicological exposure conditions for algae²⁰, and relevant to environmental exposures. 98 99 Suspensions of 100 ppb were prepared in each media and left under ambient laboratory 100 conditions for 72 hours. Aliquots were withdrawn at 0, 24, 48 and 72 hours for testing. All 101 experiments performed in triplicate. The suspension was not shaken after mixing and no 102 particle precipitation was observed.

103

104 Characterization. Measurements were performed on a Malvern Instruments Nanosizer to 105 collect both size and zeta potential data. For the size measurements 1 ml of particle suspension 106 was placed into a polystyrene disposable cuvette (Sarstedt AG &Co.). Ten consecutive 107 measurements were collected and averaged to calculate a Z average size. The results were taken at 20 °C with samples equilibrated for 2 min before measurements were started. The 108 109 Stokes relationship was used to calculate the hydrodynamic diameter of the particles. For the 110 zeta potential measurements 1 ml of suspension was put into a disposable folded capillary cell 111 (Malvern Instruments Ltd.). The sample was allowed to equilibrate at 20 °C for 2 minutes 112 before five consecutive measurements were made.

113

Flow field flow fractionation (FIFFF), high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) and electron energy loss spectroscopy (EELS) were

116 also used to examine the NPs. FIFFF separation and sizing were carried out on a Postnova 117 asymmetrical field-flow fractionation (AF2000 Mid Temperature, Postnova Analytic). The 118 accumulation wall was a 1 KDa regenerated cellulose membrane. The eluent was 0.01 M NaCl (pH 7.5). The channel flow was 1 mL min⁻¹ and cross-flow was 2 mL min⁻¹, to ensure a good 119 120 separation between the void peak and particle elution time. The injection volume was 0.5 mL 121 (particle concentration of approximately 100 ppb), which was injected into the channel after 6 122 min. The channel volume was calculated using 20, 30, and 60 nm polystyrene bead standards 123 (Duke Scientific Corp.). All particles were detected with a UV detector at 254 nm. Diffusion coefficients were calculated using FIFFF theory²¹ and converted to size using the Stokes 124 125 relationship. At least 3 replicates were collected and a mean size calculated.

126

127 Samples for HAADF-STEM analysis were prepared by placing an amorphous carbon coated copper grid (Agar Scientific, UK) onto a bespoke Teflon flat surface in a 12ml 128 ultracentrifugation tube as we have previously preformed²². 11 ml of NP suspension was 129 130 placed into the tube and centrifuged at 500,000 g for 1 hour using a Beckman ultracentrifuge 131 (L7-65 Ultracentrifuge) with a swing out rotor (SW40Ti) on to a carbon coated copper TEM 132 grid. The supernatant, containing the ionic fraction of the suspension, was removed and 133 discarded while the grid was recovered and washed by carefully placing into UHP water for 5 134 minutes. The grid was removed from the wash water and allowed to fully dry at room 135 temperature for at least two hours before imaging. The maximum time between grid 136 preparation and imaging was kept to below 12 hours.

137

138 HAADF-STEM and EELS data were obtained at 200 KeV on a Jeol 2100F coupled with a 139 CEOS spherical aberration probe corrector and a Gatan Enfina EELS. The STEM images were 140 used to analyze and compare NP morphology within the samples. The STEM probe size is 141 circa 0.1 nm. The lattice spacing for each suspension were measured using Digital Micrograph 142 software by taking line profiles across the (111) lattice direction on the HAADF-STEM 143 micrographs. Lattice spacings were measured for at least 30 NPs in each suspension. A student 144 t-test was performed between all lattice spacing sets (NPs as prepared and in media) to 145 calculate any significant changes after addition to exposure media.

146

The oxidation state of the NPs was investigated using EELs. The spatial resolution of the EELS is comparable to that of the STEM probe (0.1 nm) and the energy resolution in the order of 1 eV. Pixel sizes of 0.065 - 1 nm were used in recording one-dimensional EELS spectra, dependent on the particle size. Cerium oxidation state is sensitive to the electron beam with prolonged beam time. To minimize this damage the total time over a single point was kept to 3 seconds, as performed previously.²³ Line acquisitions took three 1 s spectra for each point along the line and the mean calculated ^{4, 24}.

154

The EELS spectra of cerium is characterized by two sharp edges at 903 eV (M4 edge) and 886 eV (M5 edge) that are due to the transition of a core electron to an unbound state, $3d3/2 \rightarrow 4f5/2$ and the $3d5/2 \rightarrow 4f7/2$, respectively (An in depth discussion of the EELS analysis can be found in a previous paper Merrifield et al 2013^4). There are many different methods employed to analyze the EELS spectra to obtain the Ce(III):Ce(IV) ratio²⁵ but here the second derivative method was used, as this has been shown to be less susceptible to alterations in sample thickness ²⁶. The integrated intensity of each edge was calculated between the points of inflection in the second derivative to find the ratio between the M5 and M4 edge. The raw EELS signal was background corrected, low pass filtered, and the second derivative calculated using the digital micrograph software. The M5/M4 ratio was then determined using the integrated intensities of the peaks in the second derivative.

166

167 The relative positions, intensities and shapes of the M4 and M5 edges present in the EELS spectra differently for the Ce(III) and Ce(IV) spectra⁴. In comparison to the Ce(III) signal the 168 169 edges in the Ce(IV) spectra are shifted to a slightly higher energy and a shoulder appears to on the M4 edge²⁷. The ratio of the edge maximum intensities also alters. The ratios of these peaks 170 171 (M5/M4) can vary somewhat in the literature and have been quoted to be anywhere between 0.79-0.91 and 1.11-1.31 for Ce(IV) and Ce(III) respectively^{4, 13, 28} showing the importance of 172 173 calibrating to a known standard. In this work, an M5/M4 ratio of 0.80 (\pm 0.1) and 1.23 (\pm 0.15) were obtained for Ce(IV) and Ce(III) standards, respectively⁴. 174

175

177 RESULTS AND DISCUSSION

178

179 The hydrodynamic diameter data (measured by DLS and FIFFF) and electrophoretic mobility 180 (EPM) of particles in exposure media at time 0 and 72 h are summarized in Table 1 (data in 181 supporting information Figures S2A and B). All of the suspensions, with the exception of one, 182 had hydrodynamic diameters of in the range 6-9 nm and there was no significant difference 183 between the as-prepared NPs and the NPs in media. However, the Ce(III) NPs in soft water 184 containing SRFA gave an immediate increase in hydrodynamic diameter to > 120 nm, which 185 was substantially different from the as-prepared size of 6.2 ± 0.1 nm. This increase was 186 significantly different from all of the other suspensions ($p \ll 0.05$). Interestingly, the FIFFF 187 measurements showed no significant differences in any of the sizes (p > 0.05), including 188 Ce(III) in soft water with SRFA. However the intensity data for these NPs was less than 50% 189 of the other conditions, indicating losses, most likely due to aggregation, as we have seen 190 before³. The combination of the two methods allowed improved identification of 191 transformation: Ce(III) NPs in soft water containing SRFA underwent some aggregation, based 192 on DLS data which is biased towards larger particles and loss of peak signal in FFF, with over 193 50% of the initial mass present in the aggregates and slightly under 50% remaining as the 194 original dispersed NPs (from intensity data). Likely this agglomeration was from a mixture of 195 the starting components: mixtures of FA, partly degraded ceria NPs and cerium ions. This 196 assumption is supported by the STEM-EELS data provided later. Changes in electrophoretic 197 mobility were also measured upon addition into the media. Certain general observations were 198 clear: Firstly Ce(IV) NPs were always negatively charged and showed no temporal changes. 199 Secondly for both NPs, the electrophoretic mobility appears to be controlled by FA if present,

as expected^{29, 30}, and showed no temporal changes, suggesting interactions were rapid 200 201 compared to the measurement time. In addition, Ce (III) NPs showed some significant 202 temporal changes in the Daphnia media and softwater, which were broadly comparable and 203 composed of only major ions, lacking the transition metals or FA present in the other media 204 (Figure S3). Previously, we have seen no effect on PVP aggregation with ionic strength,³¹ although we have seen alterations in some PVP behavior with increasing ion concentrations.³² 205 206 Speculatively, the increased ionic strength or specific ion concentration affects the polymer 207 chemistry, possibly affecting the nature of the core redox behavior.

208

209 HAADF-STEM and EELs were utilized to investigate potentially subtle changes in 210 morphology and oxidation state of synthesized Ce(III) and Ce(IV) NPs after exposure to 211 standard ecotoxicology and synthetic environmental media. Figure 1 shows a typical HAADF-212 STEM image for the Ce(III)-stock, Ce(IV)-stock, Ce(III)-softwater with fulvic acid and 213 Ce(IV)-softwater with fulvic acid after 72 h in the suspension. All the other particles in the 214 algae, daphnia and softwater media suspensions showed no observable morphological 215 differences from the stock suspensions so were not included. The Ce(III)-softwater with SRFA 216 NPs have less ordered structure than the Ce(III)-stock suspension. Although some smaller 217 particles were present (depicted in Figure 1) in the softwater medium, there were also larger, 218 unstructured materials (shown in Figure S4). These were not observed in any other 219 suspensions. The HAADF-STEM images agreed with the previously shown DLS and FIFFF 220 data, indicating that the presence of SRFA in these media resulted in alterations, including 221 formation of loosely bound agglomerates of partially degraded ceria NPs and FA. The changes 222 observed were not seen in other NPs in this study or in PVP stabilized NPs with other core

materials³¹), suggesting that media-induced transformations are dependent on core material, as well as coating and media conditions. The (111) lattice spacing for the Ce(III)-stock and Ce(IV)-stock NPs were measured as 0.34 (\pm 0.09) and 0.32 (\pm 0.01), very similar to literature data¹³). These values did not change significantly for any of any NPs in any media, with the exception of the Ce(III)-softwater with fulvic acid suspension, where no measurement was possible due to aggregation.

229

230 The oxidation state of the NPs was investigated using EELs. Both spatially averaged and 231 spatially resolved data was recorded. The M5/M4 was used to derive oxidation state ratio with reference Ce(III) and Ce(IV) measured as 1.23 (\pm 0.15) and 0.80 (\pm 0.10) respectively⁴, in 232 agreement with data in the literature ^{4, 33}. Figures in brackets are standard deviations. These 233 234 values were used to assess the oxidation state of the NPs as prepared and in the various media. 235 The mean oxidation state for each suspension was measured by comparing the ratio of the M5/M4 peaks in the EELs spectra as described in the literature⁴. As expected there is a 236 significant difference between the Ce(III)-stock and Ce(IV)-stock suspensions with p < 0.05237 238 showing that initially the particles were in different oxidation states.

239

The M5/M4 ratios for the different suspensions of originally Ce(III) after 72 h can be seen in Figure 2 (Raw data can be found in Figure S5). T-tests were performed, primarily with the aim of identifying any changes in oxidation state due to the exposure to different media types compared with the same NPs in the stock. (A table of p values can be found in Table S6). For Ce(III) NPs, significant changes occurred after spiking into the algal and daphnia exposure media, but not into the soft water either with or without SRFA. It should be noted first that the

246 Ce(III)-softwater with SRFA has a larger error related to the aggregation and amorphous nature 247 of the particles making accurate quantification difficult and, secondly, that these significant 248 differences were small in magnitude. The origin of these changes is not clear, with the primary 249 difference in the medium being the higher ionic strength of the exposure media compared to 250 the soft water. The greater complexity of the algal media containing redox active transition 251 metals plays no obvious role. However, the changes are small not significant, with the 252 exception of the dramatic effect of the SRFA on Ce(III). It is likely if there is a redox effect due to the SRFA causing oxidation and (partial) dissolution of the ceria NP, possibly 253 enhancing polymer bridging³⁴ between PVP and hence forming weakly bound agglomerates. 254 255 This interaction is in contrast to the lack of interaction with PVP on gold NPs measured using different methods³¹, suggesting a combined effect of core transformations disruption of the 256 257 PVP coating and ceria, cerium and SRFA interaction to form loosely bound agglomerates.

258

259 In the case of Ce(IV) NPs a significant (p < 0.05) and larger change in oxidation state was 260 observed upon spiking NPs into the algal and daphnia media, but there was no corresponding 261 change in EPA softwater either with or without SRFA. Semi-quantitatively, the reduction was 262 between 0-50%, dependent on new media (Figure 2; Table S7), although there was some 263 spatial variability between and within single particles. The relatively simple composition of 264 daphnia media suggests that the high ionic strength and presence of specific ions rather than 265 the presence of transition metals affect redox behavior. Most likely, the interactions of ions 266 with the PVP caused an indirect effect on the ceria core redox behavior, although the exact 267 mechanism is not clear. Taking Figures 1 and 2 together, the SRFA media reduces transformations of the Ce(IV) NPs, while the lower ionic strength media has little effect. The 268

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- diametrically opposite behavior of FA with Ce(III) and Ce(IV) is noteworthy, given thedifferent likely behaviors of these NPs to identical solution conditions.
- 271

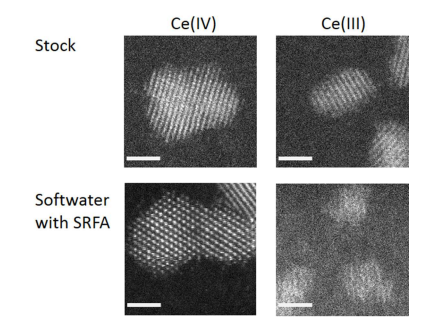
272 Examples of spatially resolved HAADF-STEM-EELS data on individual NPs are shown in 273 Figure 3. Examples for Ce(IV)-stock, Ce(IV)-algae and Ce(IV)-softwater with fulvic acid 274 micrographs are shown with corresponding M5/M4 ratios along a radial profile. In the 275 Ce(IV)-stock and Ce(IV)-softwater with fulvic acid case the M5/M4 ratio remains consistently 276 close to 0.8 from the center to the edge of the particles. Interestingly, the NPs in the SRFA 277 containing media show a more consistent pattern with no change. However, in the stock, there 278 is an indication the M5/M4 ratio increases towards the edge, showing a greater degree of Ce(III) character at the edge as would be expected³⁵. This result highlights the stability of the 279 280 Ce(IV) oxidation state in these samples, particularly in the presence of SRFA. In contrast, in 281 the Ce(IV) NPs in algal media, there is a variation in M5/M4 ratio throughout the particle. 282 These changes appear to be random, but do indicate a change from a predominantly Ce(IV) 283 oxidation state at the core of the particle to a mixed (III, IV) state. Again, there is an indication 284 that the oxidation state becomes predominantly Ce(III) at the edge. It is generally accepted that the Ce(III) oxidation state is more energetically favorable for small cerium NPs ³⁶, agreeing 285 286 with our data. In the absence of the protective SRFA, the tendency might be for these NPs to 287 become dominated by the Ce(III) state. There have been discussions in the (eco)toxicology 288 community regarding the use of dispersants and whether or not to use natural organic 289 macromolecules (NOM) as a 'natural dispersant' to overcome aggregation and other transformations³⁷. The data here suggests that, for Ce-containing NPs and perhaps other similar 290 291 NPs, NOM may be useful in controlling transformations. However, in certain cases such as the

ceria (III) NPs, NOM may also enhance transformations. In spite of this, NOM additions may
be useful since they lead to conditions which more closely resemble exposure conditions in the
environment.

295

296 The observed results have important implications for the fate and behavior, bioavailability and 297 toxicology of ceria NPs. The high resolution physical and chemical characterization of NP 298 dynamics has been essential in obtaining data on these subtle changes in NP in media relevant 299 to ecotoxicology exposures and media. In particular, exposure of NPs to inorganic standard 300 exposure media leads to small but potentially significant changes in NP oxidation state. The 301 role of NOM such as SRFA is more pronounced but causes very different transformational 302 behaviors depending on the starting NP, with Ce(III) NPs changing and agglomerating, while 303 Ce(IV) NPs appear to be stabilized and transformations minimized. This, and similar high resolution TEM-EELS studies on silver NPs^{24, 38}, need to be incorporated in future research to 304 305 understand better the full extent of these subtle transformations, which can then be linked 306 mechanistically to transport and bioavailability studies.

308 FIGURES

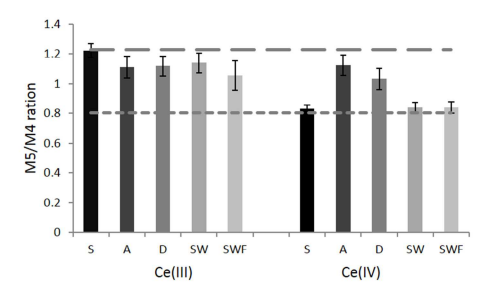


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- 310 Figure 1: Typical STEM images of Ce(IV)-stock, Ce(III)-stock, Ce(IV)-softwater with SRFA
- 311 and Ce(III)-softwater with SRFA. The bars in the image are 2 nm.

312

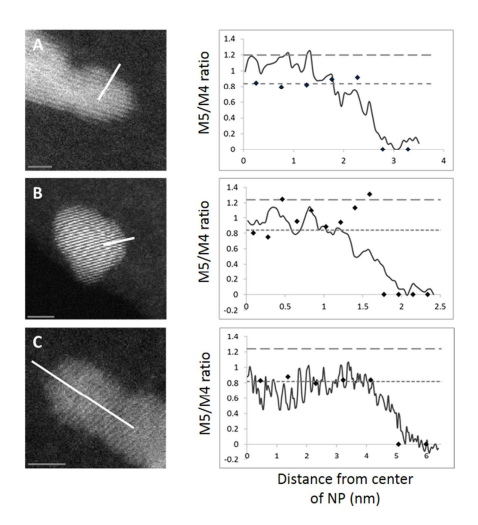
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316 Figure 2: The average ratios of the M5/ M4 cerium peaks from EELs spectrum for the centre of

the particles when exposed to different media. The dashed line is the expected ratio for pure
Ce(III) (1.23), the dotted line is the ratio for pure Ce(IV) (0.802). S - stock; A –algal media; D
– daphnia media; SW – soft water; SWF – soft water with SRF



321

Figure 3. HAADF-STEM micrographs with corresponding spatially resolved EELs analysis. Line intensity profiles (solid dark gray line) were taken from the corresponding lines (white) on the HAADF-STEM micrographs with corresponding M5/M4 ratios from the EELS analysis (black diamonds) for three suspensions A) Ce(IV)-stock, B) Ce(IV)-algae, and Ce(IV)softwater with SRFA NPs from suspension. The M5/M4 ratios for Ce(III) and Ce(IV) reference samples are shown by the light grey dashed and dotted horizontal lines respectively. Scale bars are 2 nm shown in grey on the images.

329

331 TABLES.

| | DLS | | | | Zeta | | | | FFF | | | |
|------------|--------------|------|--------------|------|---------|------|----------|------|--------------|------|--------------|------|
| 0 ho | | ours | rs 72 hou | | 0 hours | | 72 hours | | 0 hours | | 72 hours | |
| | Size (nm) | PDI | Size (nm) | PDI | | SD | | SD | Size (nm) | SD | Size (nm) | SD |
| Ce(III)S | 6.2 | 0.13 | 6.2 | 0.13 | 0.43 | 0.15 | 0.43 | 0.15 | 7.43 | 2.15 | | |
| Ce(III)A | 8.6 | 0.4 | 7.5 | 0.3 | -10.15 | 0.69 | -7.3 | 1.96 | 6.58 | 1.14 | 7.26 | 1.23 |
| Ce(III)D | 6.6 | 0.8 | 7.3 | 0.3 | 10.47 | 0.56 | -2.66 | 1.45 | 7.20 | 1.11 | 7.29 | 1.15 |
| Ce(III)SW | 7.9 | 0.5 | 7.2 | 1.0 | 8.60 | 0.63 | -6.36 | 0.33 | 6.50 | 0.84 | 6.71 | 1.26 |
| Ce(III)SWF | 122.8 | 0.3 | 122.6 | 0.3 | -14.10 | 1.13 | -15.70 | 0.14 | 8.05 | 1.15 | 7.65 | 1.08 |
| Ce(IV)S | 6.9 | 0.2 | 6.9 | 0.2 | -17.20 | 0.95 | -17.20 | 0.95 | 6.90 | 1.13 | | |
| Ce(IV)A | 7.9 | 0.5 | 7.8 | 0.4 | -23.20 | 1.27 | -19.22 | 1.71 | 7.02 | 1.16 | 7.50 | 1.17 |
| Ce(IV)D | 8.1 | 0.7 | 8.5 | 0.1 | -4.04 | 1.31 | -7.62 | 1.54 | 7.16 | 1.14 | 7.49 | 1.17 |
| Ce(IV)SW | 7.6 | 0.5 | 8.3 | 0.6 | -14.40 | 1.52 | -14.71 | 0.71 | 7.68 | 1.12 | 7.75 | 1.12 |
| Ce(IV)SWF | 8.4 | 0.7 | 8.8 | 0.8 | -12.40 | 3.15 | -14.80 | 0.99 | 7.25 | 1.04 | 7.26 | 1.06 |

Table 1. Hydrodynamic diameters (measured by FIFFF and DLS) and surface potential (measured by Zeta potential) of NP suspensions in different media at time 0 and 72 hours (Please refer to the key in S1-A for annotation names).

| 350 | Supporting information is provided on sample information, media contents, DLS results, |
|-----|--|
| 351 | FIFFF results, Z-potential results, further STEM image of Ce(III) _{SWF} NPs, Ce(III) and Ce(IV) |
| 352 | standards for EELS and original M5/M4 ratio data. This material is available free of charge via |
| 353 | the Internet at <u>http://pubs.acs.org</u> . |

- 355 AUTHOR INFORMATION
- 356 **Corresponding Author**
- 357 * Prof. Jamie R. Lead. <u>JLEAD@mailbox.sc.edu</u>.

358 Author Contributions

- 359 The manuscript was written through contributions of all authors. All authors have given
- approval to the final version of the manuscript.
- 361

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- 366 References
- 367 1. Lowry, G. V.; Gregory, K. B.; Apte, S. C.; Lead, J. R., Transformations of
- Nanomaterials in the Environment. *Environmental Science & Technology* 2012, *46*, (13), 68936899.
- 2. Romer, I.; Gavin, A. J.; White, T. A.; Merrifield, R. C.; Chipman, J. K.; Viant, M. R.;
- 371 Lead, J. R., The critical importance of defined media conditions in Daphnia magna
- 372 nanotoxicity studies. *Toxicol. Lett.* **2013**, *223*, (1), 103-108.
- 373 3. Romer, I.; White, T. A.; Baalousha, M.; Chipman, K.; Viant, M. R.; Lead, J. R.,
- Aggregation and dispersion of silver nanoparticles in exposure media for aquatic toxicity tests.
 J. Chromatogr. A 2011, *1218*, (27), 4226-4233.
- 4. Merrifield, R. C.; Wang, Z. W.; Palmer, R. E.; Lead, J. R., Synthesis and
- characterization of polyvinylpyrrolidone coated cerium oxide nanoparticles. *Environmental science & technology* 2013, *47*, (21), 12426-33.
- 379 5. Graham, U. M.; Tseng, M. T.; Jasinski, J. B.; Yokel, R. A.; Unrine, J. M.; Davis, B. H.;
- 380 Dozier, A. K.; Hardas, S. S.; Sultana, R.; Grulke, E. A.; Butterfield, D. A., In Vivo Processing
- 381 of Ceria Nanoparticles inside Liver: Impact on Free-Radical Scavenging Activity and
- 382 Oxidative Stress. *Chempluschem* **2014**, *79*, (8), 1083-1088.
- 383 6. Zantye, P. B.; Kumar, A.; Sikder, A. K., Chemical mechanical planarization for
 384 microelectronics applications. *Materials Science & Engineering R-Reports* 2004, 45, (3-6), 89385 220.
- 386 7. Cumbo, M. J.; Fairhurst, D.; Jacobs, S. D.; Puchebner, B. E., Slurry particle size
- evolution during the polishing of optical glass. *Appl. Opt.* **1995**, *34*, (19), 3743-3755.
- 8. Hoshino, T.; Kurata, Y.; Terasaki, Y.; Susa, K., Mechanism of polishing of SiO2 films
 by CeO2 particles. *Journal of Non-Crystalline Solids* 2001, *283*, (1-3), 129-136.
- Stanek, C. R.; Tan, A. H. H.; Owens, S. L.; Grimes, R. W., Atomistic simulation of
 CeO(2) surface hydroxylation: implications for glass polishing. *Journal of Materials Science*
- **2008**, *43*, (12), 4157-4162.
- 393 10. Sajith, V.; Sobhan, C. B.; Peterson, G. P., Experimental Investigations on the Effects of
 394 Cerium Oxide Nanoparticle Fuel Additives on Biodiesel. *Advances in Mechanical Engineering*395 2010.
- Johnson, A. C.; Park, B., Predicting contamination by the fuel additive cerium oxide
 engineered nanoparticles within the United Kingdom and the associated risks. *Environmental Toxicology and Chemistry* 2012, *31*, (11), 2582-2587.
- 399 12. Karakoti, A. S.; Munusamy, P.; Hostetler, K.; Kodali, V.; Kuchibhatla, S.; Orr, G.;
- 400 Pounds, J. G.; Teeguarden, J. G.; Thrall, B. D.; Baer, D. R., Preparation and characterization
- 401 challenges to understanding environmental and biological impacts of ceria nanoparticles.
- 402 Surface and Interface Analysis **2012**, *44*, (8), 882-889.
- 403 13. Deshpande, S.; Patil, S.; Kuchibhatla, S.; Seal, S., Size dependency variation in lattice
 404 parameter and valency states in nanocrystalline cerium oxide. *Applied Physics Letters* 2005,
 405 87, (13).
- 406 14. Ohta, A.; Kawabe, I., REE(III) adsorption onto Mn dioxide (δ -MnO2) and Fe
- 407 oxyhydroxide: Ce(III) oxidation by δ -MnO2. Geochimica et Cosmochimica Acta 2001, 65, (5),
- 408 695-703.

- 409 15. Xue, Y.; Zhai, Y. W.; Zhou, K. B.; Wang, L.; Tan, H. N.; Luan, Q. F.; Yao, X., The
- Vital Role of Buffer Anions in the Antioxidant Activity of CeO2 Nanoparticles. *Chemistry-a European Journal* 2012, *18*, (35), 11115-11122.
- 412 16. Shah, V.; Shah, S.; Shah, H.; Rispoli, F. J.; McDonnell, K. T.; Workeneh, S.; Karakoti,
- 413 A.; Kumar, A.; Seal, S., Antibacterial Activity of Polymer Coated Cerium Oxide
- 414 Nanoparticles. *Plos One* **2012**, *7*, (10).
- 415 17. Collin, B. A., Melanie;; Johnson, Andrew; Kaur, Inder; Keller, Arturo; Lazareva,
- 416 Anastasiya; Lead, Jamie; Ma, Xingmao; Merrifield, Ruth; Svendsen, Claus; White, Jason;
- 417 Unrine, Jason., Environmental release, fate and ecotoxicological effects of manufactured ceria
- 418 nanomaterials. *Environmental Science: Nano* 2014, Accepted
- 419 18. Auffan, M.; Rose, J.; Orsiere, T.; De Meo, M.; Thill, A.; Zeyons, O.; Proux, O.;
- 420 Masion, A.; Chaurand, P.; Spalla, O.; Botta, A.; Wiesner, M. R.; Bottero, J.-Y., CeO2
- 421 nanoparticles induce DNA damage towards human dermal fibroblasts in vitro. *Nanotoxicology*422 **2009**, *3*, (2), 161-U115.
- 423 19. Peng, L.; He, X.; Zhang, P.; Zhang, J.; Li, Y. Y.; Zhang, J. Z.; Ma, Y. H.; Ding, Y. Y.;
- 424 Wu, Z. Q.; Chai, Z. F.; Zhang, Z. Y., Comparative Pulmonary Toxicity of Two Ceria
- 425 Nanoparticles with the Same Primary Size. International Journal of Molecular Sciences 2014,
- 426 15, (4), 6072-6085.
- 427 20. OECD GUIDELINES FOR THE TESTING OF CHEMICALS PROPOSAL FOR
- 428 UPDATING GUIDELINE 201: Freshwater Alga and Cyanobacteria, Growth Inhibition Test.
 429 *http://www.oecd.org/chemicalsafety/testing/1946914.pdf* 2011.
- 430 21. Baalousha, M.; Stolpe, B.; Lead, J. R., Flow field-flow fractionation for the analysis 431 and characterization of natural colloids and manufactured nanoparticles in environmental
- 432 systems: A critical review. J. Chromatogr. A **2011**, *1218*, (27), 4078-4103.
- 433 22. Prasad, A.; Lead, J. R.; Baalousha, M., An electron microscopy based method for the
- detection and quantification of nanomaterial number concentration in environmentally relevant
 media. *Science of the Total Environment* 2015, *537*, 479-486.
- 436 23. Baalousha, M.; Ju-Nam, Y.; Cole, P. A.; Hriljac, J. A.; Jones, I. P.; Tyler, C. R.; Stone,
- 437 V.; Fernandes, T. F.; Jepson, M. A.; Lead, J. R., Characterization of cerium oxide
- 438 nanoparticles-Part 2: Nonsize measurements. *Environmental Toxicology and Chemistry* 2012,
- 439 *31*, (5), 994-1003.
- 440 24. Römer, I.; Wang, Z. W.; Merrifield, R. C.; Palmer, R. E.; Lead, J., High Resolution
- 441 STEM-EELS Study of Silver Nanoparticles Exposed to Light and Humic Substances.
- 442 Environmental Science & Technology **2016**, *50*, (5), 2183-2190.
- 443 25. Manoubi, T.; Colliex, C.; Rez, P., QUANTITATIVE ELECTRON-ENERGY LOSS
- 444 SPECTROSCOPY ON M45 EDGES IN RARE-EARTH-OXIDES. Journal of Electron
- 445 Spectroscopy and Related Phenomena **1990**, *50*, (1-2), 1-18.
- 446 26. Fortner, J. A.; Buck, E. C.; Ellison, A. J. G.; Bates, J. K., EELS analysis of redox in 447 glasses for plutonium immobilization. *Ultramicroscopy* **1997**, *67*, (1-4), 77-81.
- 448 27. Turner, S.; Lazar, S.; Freitag, B.; Egoavil, R.; Verbeeck, J.; Put, S.; Strauven, Y.; Van
- 449 Tendeloo, G., High resolution mapping of surface reduction in ceria nanoparticles. *Nanoscale*
- **2011**, *3*, (8), 3385-3390.
- 451 28. Yang, G.; Möbus, G.; Hand, R., Fine structure EELS analysis of glasses and glass
- 452 composites. Journal of Physics: Conference Series 2006, 26, (1), 73.
- 453 29. Fabrega, J.; Zhang, R.; Renshaw, J. C.; Liu, W.-T.; Lead, J. R., Impact of silver
- 454 nanoparticles on natural marine biofilm bacteria. *Chemosphere* **2011**, *85*, (6), 961-966.

455 30. Osborne, O. J.; Johnston, B. D.; Moger, J.; Baalousha, M.; Lead, J. R.; Kudoh, T.; 456 Tyler, C. R., Effects of particle size and coating on nanoscale Ag and TiO2 exposure in 457 zebrafish (Danio rerio) embryos. Nanotoxicology 2013, 7, (8), 1315-1324. 458 31. Hitchman, A.; Smith, G. H. S.; Ju-Nam, Y.; Sterling, M.; Lead, J. R., The effect of 459 environmentally relevant conditions on PVP stabilised gold nanoparticles. Chemosphere 2013, 460 90, (2), 410-416. 461 32. Mirshahghassemi, S.; Cai, B.; Lead, J. R., Evaluation of polymer-coated magnetic 462 nanoparticles for oil separation under environmentally relevant conditions: effect of ionic 463 strength and natural organic macromolecules. Environmental Science-Nano 2016, 3, (4), 780-464 787. 465 Wu, L. J.; Wiesmann, H. J.; Moodenbaugh, A. R.; Klie, R. F.; Zhu, Y. M.; Welch, D. 33. O.; Suenaga, M., Oxidation state and lattice expansion of CeO2-x nanoparticles as a function 466 467 of particle size. *Physical Review B* 2004, 69, (12). 468 34. Cumberland, S. A.; Lead, J. R., Particle size distributions of silver nanoparticles at environmentally relevant conditions. J. Chromatogr. A 2009, 1216, (52), 9099-9105. 469 470 Dogra, Y.; Arkill, K. P.; Elgy, C.; Stolpe, B.; Lead, J.; Valsami-Jones, E.; Tyler, C. R.; 35. 471 Galloway, T. S., Cerium oxide nanoparticles induce oxidative stress in the sediment-dwelling 472 amphipod Corophium volutator. Nanotoxicology 2016, 10, (4), 480-487. 473 Quik, J. T. K.; Lynch, I.; Hoecke, K. V.; Miermans, C. J. H.; Schamphelaere, K. A. C. 36. 474 D.; Janssen, C. R.; Dawson, K. A.; Stuart, M. A. C.; Meent, D. V. D., Effect of natural organic 475 matter on cerium dioxide nanoparticles settling in model fresh water. *Chemosphere* 2010, 81, 476 (6), 711-715.

- 477 37. Petersen, E. J.; Diamond, S. A.; Kennedy, A. J.; Goss, G. G.; Ho, K.; Lead, J.; Hanna,
- 478 S. K.; Hartmann, N. B.; Hund-Rinke, K.; Mader, B.; Manier, N.; Pandard, P.; Salinas, E. R.;
- 479 Sayre, P., Adapting OECD Aquatic Toxicity Tests for Use with Manufactured Nanomaterials:
- Key Issues and Consensus Recommendations. *Environmental Science & Technology* 2015, *49*,
 (16), 9532-9547.
- 482 38. Baalousha, M.; Arkill, K. P.; Romer, I.; Palmer, R. E.; Lead, J. R., Transformations of
- 483 citrate and Tween coated silver nanoparticles reacted with Na2S. *Science of the Total*
- 484 *Environment* **2015**, *502*, 344-353.