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Coupled redox transformations of catechol and cerium at the surface of a cerium(III) phosphate mineral

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Abstract

Highly insoluble Ce-bearing phosphate minerals form by weathering of apatite [Ca$_5$(PO$_4$)$_3$.OH.F.CI)], and are important phosphorous repositories in soils. Although these phases can be dissolved via biologically-mediated pathways, the dissolution mechanisms are poorly understood. In this paper we report spectroscopic evidence to support coupling of redox transformations of organic carbon and cerium during the reaction of rhabdophane (CePO$_4$.C$_1$H$_2$O) and catechol, a ubiquitous biogenic compound, at pH 5. Results show that the oxic–anoxic conditions influence the mineral dissolution behavior. Under anoxic conditions, the release of P and Ce occurs stoichiometrically. In contrast, under oxic conditions, the mineral dissolution behavior is incongruent, with dissolving Ce$^{3+}$ ions oxidizing to CeO$_2$. Reaction product analysis shows the formation of CO$_2$, polymeric C, and oxalate and malate. The presence of more complex forms of organic carbon was also confirmed. Near edge X-ray absorption fine structure spectroscopy measurements at Ce-M$_{4,5}$ and C-K absorption edges on reacted CePO$_4$.C$_1$H$_2$O samples in the absence or presence of catechol and dissolved oxygen confirm that (1) the mineral surface converts to the oxide during this reaction, while full oxidation is limited to the near-surface region only; (2) the Ce valence remains unchanged when the reaction between CePO$_4$.H$_2$O and O$_2$ but in the absence of catechol. Carbon K-edge spectra acquired from rhabdophane reacted with catechol under oxic conditions show spectral features before and after reaction that are considerably different from catechol, indicating the formation of more complex organic molecules. Decreases in intensity of characteristic catechol peaks are accompanied by the appearance of new $\pi^*$ resonances due to carbon in carboxyl (ca. 288.5 eV) and carbonyl (ca. 289.3 eV) groups, and the development of broad structure in the $\pi^*$ region characteristic of aliphatic carbon. Evolution of the C K-edge spectra is consistent with aromatic-ring cleavage and polymerization. These results further substantiate that the presence of catechol, O$_2$(aq) causes both the oxidation of structural Ce$^{3+}$ and the transformation of catechol to more complex organic molecules. Scanning Transmission X-Ray Microscopy measurements at the C K and Ce M$_{4,5}$ edges indicate three dominant organic species, varying in complexity and association with the inorganic phase. Untransformed catechol is loosely associated with CeO$_2$, whereas more complex organic molecules that exhibit lower aromaticity and stronger C=O $\pi^*$ resonances of carboxyl-C and carbonyl-C groups are only found in association with the grains. These results further serve as basis to postulate that, in the presence of O$_2$, CeO$_2$ can mediate the oxidative polymerization of catechol to form higher molecular weight polymers. The present work provides evidence for a pathway of biologically-induced, non-enzymatic oxidation of cerium and formation of small CeO$_2$ particles at room temperature. These findings may have implications for carbon cycling in natural and cerium-contaminated soils and aqueous environments.

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1. INTRODUCTION

Phosphate bioavailability can limit ecosystem productivity. In near-surface low-temperature geological settings, phosphorus is generally partitioned between the aqueous phase, organic matter, mineral surfaces, primary phosphate phases (such as apatite) and secondary phosphates (commonly trivalent metal phosphate minerals) (Banfield and Eggleton, 1989). For microbial productivity, phosphate limitation may occur due to sequestration of phosphate into insoluble secondary phosphate minerals or through incorporation of phosphate into recalcitrant organic complexes, such as phosphohumate complexes (Crews et al., 1995). The formation of secondary phosphate phases is a function of the localized availability of phosphate and lanthanide ions. For example, microcrystalline lanthanide-bearing phosphate minerals may form as a result of apatite \([\text{Ca}_5(\text{PO}_4)_3(\text{OH},\text{Cl,F})]\) and allanite \([\text{Ce}_2(\text{Al})_3(\text{SiO})_4\text{OH}]\) weathering.

Microcrystalline lanthanide-bearing phosphate minerals are important metal and phosphorous reservoirs in soils and other subsurface environments (Banfield and Eggleton, 1989). However, in soils where secondary phosphate minerals, including lanthanide phosphates, are the dominant source of phosphorous, the colonization of mineral surfaces by native microbial communities can lead to their dissolution (Taunton et al., 2000a,b). Although lanthanide phosphate minerals are highly insoluble (solubility product \(10^{-24} - 10^{-26}\); Jonasson et al., 1985), colonization of their surfaces by native microbial communities can lead to mineral dissolution (Taunton et al., 2000a,b). This has been attributed to the combined effects of the complexation of aqueous lanthanide ions by organic ligands and microbial phosphorus uptake (Taunton et al., 2000a). In some cases, dissolution of lanthanide phosphate minerals in oxidized soils is accompanied by the precipitation of nanocrystalline CeO_2 (Green et al., 2002).

Cerium is the most abundant lanthanide and generally the only one to undergo redox reactions at the Earth’s surface. In young weathering profiles, field data suggest that most Ce exists as Ce\(^{3+}\), whereas in older weathered profiles Ce\(^{3+}\) is oxidized to Ce\(^{4+}\) (Taunton et al., 2000a,b). It has been shown previously that dissolution of phosphate minerals containing Ce\(^{3+}\) is accompanied by cerium oxidation and precipitation of nanocrystalline CeO_2 (Gardner et al., 1983; Jonasson et al., 1985; Banfield et al., 1999; Taunton et al., 2000a,b; Cervini-Silva et al., 2005). These processes are believed to be responsible for the accumulation of Ce at the top of the saprolite layers of lateritic soil profiles (Braun et al., 1990) and the depletion of dissolved Ce in the oceans relative to other lanthanides (Moffett, 1990, 1993). Oxidation of Ce\(^{3+}\) by O_2 in seawater is extremely slow, but is accelerated by microorganisms (Moffett, 1990, 1993), and it has been proposed that Ce\(^{3+}\) oxidation occurs enzymatically (Moffett, 1990, 1993).

Although redox reactions involving Ce-rich phosphate minerals may affect the stability of lanthanide phosphate minerals in soil and marine environments, little attention has been paid to the role of biogenic molecules (other than enzymes) in Ce\(^{3+}\) redox transformations, or to the possible implications of such reactions for carbon cycling. Organic compounds secreted into the subsurface environment by microbes and plants can interact with surface-bound phosphate, increasing availability of phosphorus for plant uptake (Barber and Martin, 1976; Brewster et al., 1976; Nye, 1977; Kucey et al., 1989). For example, low-molecular weight organic acids can readily induce the release of phosphate through the dissolution of apatite (Welch et al., 2002). Other organics such as polysaccharides may promote mineral dissolution by complexing with ions in solution, thereby lowering the minerals’ solution saturation state (Welch and Vandevivere, 1994; Banfield et al., 1999; Welch et al., 1999). In addition, biogenic compounds such as citrate, secreted by some plants, may form polymers with iron and phosphate, which can effectively increase their concentration in the soil solution in proximity to the plant’s roots (Gardner et al., 1983). For example, the secretion of citrate by *Lupinus albus* L. \((\leq 188\ mg\ citrate\ per\ total\ weight\ (g)\ of\ soil\ at\ pH\ 5)\) causes the proximal dissolution of phosphate ferric hydroxides and the concomitant release of iron and phosphate. Dissolved Fe and citrate then form polymers such as \([\text{FeO}_{0.95}(\text{OH})_{0.75}(\text{citrate})_{0.15}\text{Na}_{0.24}]_n\) (Gardner et al., 1983).

Abiotic experimentation provides evidence for the significance of biogenic compounds in mineral dissolution. For instance, small organics such as malate (*Jauregui and Reisenauer*, 1982), glutamate (*Traina and Doner*, 1985), diphenolics, salicylic acid, pyruvic acid, oxalic acid, fulvic acid (*Stone and Morgan*, 1984, 1987), or even monophenolic compounds sharing electron donating substituents (*Lehmann et al.*, 1987; *Stone, 1987; McBride, 1989a,b) facilitate the reductive dissolution of Mn oxides and become oxidized as a result. Because Ce\(^{4+}\) is a stronger oxidant than manganese (*Larson and Hufnal*, 1980; *Machjer et al.*, 2000), it is likely to participate in similar geochemical cycles to Mn\(^{4+}\).

We investigate here the redox mineral transformations during the reaction of CePO_4 \(\cdot\) H_2O with catechol \((C_6H_4O_2)\). Catechol is a ubiquitous biogenic compound known to undergo mineral-mediated electron transfer processes (*Larson and Hufnal*, 1980; *Bennett*, 1991; Welch and Ullman, 1992; *Wang and van Capellen*, 1996; *Larson*, 1998; *Kinrade et al.*, 1999; *Machjer et al.*, 2000; *Sparks*, 2003). Catechol is also typical intermediate in aerobic hydrocarbon degradation and has been shown to accelerate the weathering of primary silicates via the formation of stable complexes with Si and Al (*Bennett*, 1991; Welch and Ullman, 1992; *Wang and van Capellen*, 1996; *Kinrade et al.*, 1999). We show that catechol induces oxidation of Ce\(^{3+}\) to form CeO_2 and is degraded by CeO_2 production at room temperature. The outcome of this work contributes to a better understanding of the effect of biological activity on the stability of secondary phosphate minerals and their transformation is relevant to predictions of the stability of lanthanide phosphate compounds. In this paper, we aim to study rare-earth orthophosphates such as rhabdophane (CePO_4 \(\cdot\) H_2O) are amongst the most insoluble materials (solubility product \(K_{sp} = 10^{-22}\); Firschberg and Brune, 1991). Due to their low solubility, they are used as chemical tracers of natural fluid–rock processes (*Ushakov et al.*, 2001), have been studied extensively for \(c\) long-term storage.
of nuclear wastes, and are under consideration as host phases for actinides.

2. MATERIALS AND METHODS

Materials. Catechol (99% powder; CAS Number: 120-80-9), oxalic acid (99% ≤; CAS Number: 144-62-7), Malic acid (CAS Number: 6915-15-7), humic acid (CAS Number: 1415-93-6), and cerium oxide (CAS Number: 1306-38-3) were purchased from Sigma–Aldrich. Natural humic samples were kindly arranged by Prof. Garrison Sposito (University of California) and referred thereafter as compounds 1, 2, and 3 (peroxidase extracted humic acid), 4 (4,5-dichlorocatechol), 5 (2,3,4-trihydroxybenzoic acid), and 6 (Greenfield humic acid).

2.1. Mineral characterization

Cerium phosphate was synthesized by rapidly mixing 0.5 M phosphoric acid and cerium chloride solutions at 20 °C, as described elsewhere (Hikichi et al., 1978). Solid samples were separated from the liquor and rinsed with distilled water until the pH of the supernatant solution was 5.7 ± 0.2. pH measurements were conducted before conducting dissolution experiments.

Powder X-ray diffraction (XRD) analysis confirmed the solid to be a mixture of rhabdophane and monazite, polymorph phases of CePO\(_4\). The specific surface areas of the solid to be a mixture of rhabdophane and monazite, conducting dissolution experiments. 

2.2. Mineral dissolution

Dissolution of CePO\(_4\)H\(_2\)O was investigated in batch mode in filter-sterilized (0.2 μm filter) 1 mM catechol at pH 5.0, solids concentration of 1 g L\(^{-1}\), constant stirring, under oxic and anoxic conditions for periods of 7 days. Dissolution experiments under strict anoxic conditions were prepared inside a Coy environmental chamber (Grass Lake, MI) equipped with a gas analyzer and a catalytic regenerator. Anoxic stock solutions were prepared by degassing distilled water with high purity nitrogen gas (15 min L\(^{-1}\) water). The samples were then handled in a glove box. The content of dissolved oxygen during the experiments was monitored using an oxygen meter (Unisense, Denmark). Dissolved oxygen contents measured in oxic and anoxic conditions corresponded to 8.1 ± 0.1 and ≤0.01 mg L\(^{-1}\), respectively. All batch experiments conducted under anoxic conditions were septum sealed and covered with parafilm, and opened prior analysis inside the environmental chamber.

Aliquots of HCl or NaOH were added for pH adjustment. Variations in pH in suspensions bearing CePO\(_4\)H\(_2\)O and catechol were kept at <0.1 pH units. After equilibration, the suspensions were centrifuged and the supernatant solutions filtered for chemical analysis using 0.02 μM-Millipore syringe filters. Ce and P determinations were conducted using an inductively-coupled plasma atomic emission spectrometer (ICP-AES), with emission lines at 418.6 and 178.7 nm, respectively.

Two-hundred hour adsorption experiments were also conducted and the supernatant solutions filtered for catechol determination. Catechol ligand disappearance was quantified using a UV spectrometer at \(\lambda = 275\) nm (Larson and Hufnall, 1980). Catechol solutions used in experiments and inorganic oxidation controls were monitored spectrophotometrically.

Additional experiments were conducted to study whether the formation of CeO\(_2\) could occur in the absence of CePO\(_4\)H\(_2\)O as Ce\(^{3+}\) source. The initial reaction conditions were (a) \(5 \times 10^{-6} \) M CeCl\(_3\) and 9.9 \(\times 10^{-4}\) M catechol, (b) \(4 \times 10^{-5} \) M CeCl\(_3\) and 9.1 \(\times 10^{-4}\) M catechol, and (c) \(1.1 \times 10^{-4} \) M CeCl\(_3\) and 1 \(\times 10^{-3}\) M catechol. Determinations of [Ce\(^{3+}\)] (as total, dissolved Ce) and dissolved oxygen were conducted as described above by ICP-AES and oxygen meter, respectively. In addition, equivalent experiments investigated the ability of pure CeO\(_2\) to catalyze the oxidation of catechol in the presence of O\(_2\).

2.3. Reaction products analysis

2.3.1. Bulk solution and head space

Identification and quantification of reaction products in supernatant solutions was conducted using a Shimadzu SPD high pressure liquid chromatograph equipped with a UV detector and a Hamilton HC-75 column (protonated form). A 20-min program was used with a flow rate of 0.4 mL min\(^{-1}\) and a mobile phase of 0.016 N H\(_2\)SO\(_4\) in distilled water. Elution times for oxalate, malate, and catechol were 2–3.1, 6.1–7.0, and 8.5–9.5 min, respectively.

Total organic and inorganic carbon analysis of supernatant solutions was conducted using a 1010 O-I-Analytical total organic carbon analyzer equipped with a 1051 vial multisampler (College Station, TX). Evolution of CO\(_2\) was determined by head-space sampling and analyzed using a gas chromatograph equipped with a total carbon detector 8610 SRI (Torrance, CA), and corrected for CO\(_2\) atmospheric levels at 25°C. Suspensions were not purged for CO\(_2\) trapping. An gas-tight syringe was used for headspace sampling at various time intervals. Experiments were reproduced such that vials were used for analyses only once. Experiments were conducted in 3-mL gas-tight vials.

Pentane-extractions of mineral samples in aqueous solution were conducted to recover the sorbed, non-polymeric catechol. Polymeric catechol was estimated from determinations of the carbon content in mineral samples as determined by thermogravimetry.

Reported percentage weight for the conversion of C (% wt C) in suspension are referred to initial concentrations of catechol in solution.

2.3.2. Powder X-ray diffraction

XRD analysis of the initial and transformed CePO\(_4\)H\(_2\)O material was performed by drying the wet powder on a low background quartz plate for measurement with a X’Pert Pro X-ray Diffractometer (Panalytical), using Co K(α) radiation (1.7903 Å).
2.3.3. X-ray spectroscopy and spectromicroscopy

Near edge X-ray absorption fine structure (NEXAFS) spectroscopy measurements at Ce M\textsubscript{4.5} and C K absorption edges on weathered CePO\textsubscript{4}H\textsubscript{2}O samples in the absence or presence of catechol and dissolved oxygen were conducted on beamline 4.0.2 at the Advanced Light Source, Lawrence Berkeley National Laboratory (Young et al., 2001). The spectroscopic data were obtained by scanning the X-ray energy through the Ce M\textsubscript{4.5} edges with an approximate energy resolution of 0.2 eV and measuring the absorption signal through the total electron yield (TEY) with a channeltron electron-multiplier. The raw TEY signal was normalized by the incident beam intensity (I\textsubscript{0}), a smooth polynomial background and an arctangent step function were fitted to the data and the absorption step was set to unity.

Scanning transmission X-ray microscopy (STXM) measurements at the C K and Ce M\textsubscript{4.5} edges were conducted at the Molecular Environmental Science beamline 11.0.2 of the Advanced Light Source at Lawrence Berkeley National Laboratory (Tyliszczak et al., 2004). The STXM uses a Fresnel zone plate lens to focus a monochromatic soft X-ray beam onto a sample that is X-Y raster scanned. Transmitted photons were detected via a phosphor scintillator photomultiplier tube assembly, linear up to 15 MHz. This technique, pioneered by Kirz et al. (1995), is based on core electron excitation by X-ray absorption, which provides the imaging contrast. Transmission images at energies below and at the relevant absorption edges and subsequently converted into optical density images were used to derive chemical maps. Image sequences (called stacks) acquired at multiple energies spanning the relevant element absorption edge were used to extract NEXAFS spectra from areas of interest on the samples. The theoretical spectral resolution during our measurements was 100 meV and the energy was calibrated at the C K-edge using the 3p Rydberg peak of gaseous CO\textsubscript{2} at 294.96 eV, and at the Ce M\textsubscript{4.5}-edges using the Ce M\textsubscript{5}-edge of CePO\textsubscript{4}H\textsubscript{2}O at 882.8 eV.

To check for radiation induced damage on our organic samples, we collected fast stacks, using fewer spectral and spatial points and fast dwell times (0.2 ms) using small slit sizes in regions similar to the ones of interest and compared them with finer stacks performed on regions of interest. No alteration of the spectra was observed under careful conditions (incident flux <10\textsuperscript{7} photons/s, fast dwell times <0.5 ms). The images in each sequence were aligned via a spatial cross-correlation analysis using the aXis2000 software. Absorption spectra were extracted from groups of pixels from relevant spatial areas of the stack. Clean areas of the silicon nitride membrane were used to normalize the absorption signal obtained from the areas of interest.

One microliter of cerium weathered solutions was deposited using a micropipette onto a Si\textsubscript{3}N\textsubscript{4} window (Silson Ltd.), 100 nm thick, then air-dried. All reference compounds were in powder form and finely ground using an agate mortar and pestle. They were then suspended in MQ water, deposited onto a Si\textsubscript{3}N\textsubscript{4} membrane using a micropipette and air-dried. All samples were analyzed under He near atmospheric pressure.

3. RESULTS AND DISCUSSION

3.1. Cerium redox transformation

Fig. 1 shows the solubilization of Ce and phosphate during the batch dissolution of microcrystalline CePO\textsubscript{4}H\textsubscript{2}O in the presence of 1 mM catechol under oxic and anoxic conditions at pH 5. Concentrations of [Ce] and [P] following mineral dissolution in the presence of water only at pH\textsubscript{0} = 5 were found to be below detection limits. When CePO\textsubscript{4}H\textsubscript{2}O was dissolved in the absence of O\textsubscript{2}, Ce ions accumulated in solution (ca. 20 x 10\textsuperscript{-6} M). However, when dissolution occurred in the presence of O\textsubscript{2}, Ce concentration in solution did not surpass 6 x 10\textsuperscript{-6} M, indicating the formation of a Ce-sequestering solid product. Our results show that Ce:P molar ratios in solution after reacting CePO\textsubscript{4}H\textsubscript{2}O under anoxic and oxic conditions were found to be ca. 1:1 and 10±, respectively. In contrast, concentrations of Ce were found to be up to four times or higher for the case when experiments were conducted under anoxic conditions.

Experiments were conducted to study whether Ce oxidation could occur using another Ce(III) source. The fate of aqueous Ce(III) (as CeCl\textsubscript{3}) in the presence of O\textsubscript{2} and catechol at room temperature and pH 5 was studied. Decreases in Ce(III) concentration in solution, [Ce(III)], were determined using a UV spectrometer. Results depicted in Fig. 2 show that the reaction between aqueous Ce(III)

![Fig. 1](image-url)
and O₂ using CeCl₃ as the Ce(III) source is very slow, except in the presence of catechol. The oxidation of Ce(III) is fastest when the concentration of aqueous cerium is low ([Ce]₀ ca. 5 × 10⁻⁶ or 4 × 10⁻⁵ M) and the Ce(III)-to-catechol molar ratio is <0.05. Under these conditions, the complexion by catechol appears to mediate the oxidation of Ce(III) ions. Once formed, Ce(IV) can react with catechol to form low-molecular weight organic compounds (see below).

Cerium oxidation is inhibited at higher initial concentrations of aqueous cerium (1.1 × 10⁻⁴ M) and catechol (10⁻³ M) under oxic conditions, possibly due to the formation of Ce⁴⁺-catechol complexes such as [Ce(cat)₄]⁻⁴. Such complexes owe their existence and stability to the coordination ability of catechol (Sofen et al., 1979). It is possible that similar pathways prevail in concentrated suspensions of CePO₄·H₂O and catechol that could preclude the formation of Ce(IV)-bearing clusters (Sofen et al., 1979).

3.2. Solid product analysis

3.2.1. The formation of CeO₂

Two conventional analysis methods indicated that CeO₂ is the mineral product of the oxidation of CePO₄·H₂O in the presence of catechol. Powder X-ray diffraction patterns of synthetic CePO₄·H₂O dissolved in catechol under oxidizing conditions for several weeks showed a characteristic peak that can be assigned to the (111) reflection of CeO₂ (Fig. 3). However, the extent of transformation was small, and a full CeO₂ pattern could not be resolved. Prior work report infrared spectra of reacted CePO₄·H₂O samples indicate the presence of a doublet at 2360 cm⁻¹, characteristic of CeO₂ (Cervini-Silva et al., 2005). The oxidation of structural Ce was further confirmed by Ce NEXAFS Spectroscopy.

3.2.2. Ce and C NEXAFS spectroscopy: dissolved oxygen mediates C and Ce oxidation

Soft X-ray absorption spectroscopy at the Ce M₄,5-edges probes electronic transitions between core 3d electrons and valence 4f states, and hence is extremely sensitive to Ce valence and local bonding environment. Fig. 4a shows Ce 3d NEXAFS spectra for CePO₄·H₂O samples before and after reaction with catechol under oxic conditions, and a spectrum from CeO₂ reference powder. It is evident that the mineral surface has been extensively converted to the oxide during reaction. The data in Fig. 4a were acquired using the surface sensitive total electron yield signal. The simultaneous fluorescence yield signal (not shown) did not give good agreement between weathered CePO₄·H₂O and CeO₂, indicating that full oxidation was limited to the near surface region only. NEXAFS analysis of control samples showed that the Ce valence was unchanged following reaction with O₂ but in the absence of catechol and that no transformation of catechol is observed in the absence of O₂ (aq) (Fig. 4b).

NEXAFS spectroscopy at the C K-edge probes molecular bonding via 1s transitions to unoccupied hybridized 2p states (Stöhr, 1992). Carbon NEXAFS is highly sensitive to the presence of unsaturated molecules (double or triple bonds, π* orbitals) and saturated molecules (single bond, σ* orbitals) and different functional groups frequently produce distinct and characteristic peaks that can be used as fingerprints (Stöhr, 1992). Fig. 5 summarizes STXM derived C K-edge NEXAFS spectra of reference organic acids, acquired to assist the characterization of the organic products of reaction, including (a) the spectra of simpler organic acids and (b) the spectra of soil-derived model hemic acid compounds and catechol derivatives.

Fig. 6 shows TEY mode C 1s spectra of catechol before and after reaction with CePO₄·H₂O. The catechol C K-edge spectrum is straightforward to interpret, with strong peaks in the π* region that are attributable to aromatic and phenolic carbon (Cody et al., 1998; Scheinost et al., 2001; Schäfer et al., 2003). The spectral features for catechol after reaction are considerably different, however, indicating the formation of more complex organic molecules. The decrease in the intensity of characteristic catechol peaks is accompanied by the appearance of new π* resonances due to carbon in carboxyl (ca. 288.5 eV) and carbonyl (ca. 289.3 eV) groups, and the development of broad structure in the π* region characteristic of aliphatic carbon (Myneni, 2003). In summary, the NEXAFS spectroscopic analysis show that, in the presence of catechol, O₂ (aq) causes both the oxidation of structural Ce(III) and the transformation of catechol to more complex organic molecules.

3.3. Catechol transformation with CeO₂

STXM measurements on CeO₂ samples reacted with catechol are presented in Fig. 7. Spectra characteristic of each species present were extracted and used to derive qualitative component maps using singular value decomposition (SVD) (Koprinarov et al., 2002). SVD is an image sequence fitting procedure that computes a least squares fit of the intensity at each pixel to a linear combination of reference
spectra for each component and was performed using aXis2000 software. Three dominant organic species were found, varying in complexity and association with the inorganic phase. Untransformed catechol is loosely associated with CeO$_2$. On the other hand, more complex organic molecules that exhibit lower aromaticity and stronger C=O $\pi^*$ resonances of carboxyl-C and carbonyl-C groups are only preferentially found in association with the grains. These results show that, in the presence of O$_2$, CeO$_2$ can mediate the oxidative polymerization of catechol to form higher molecular weight polymers.

### 3.4. Catechol transformation with CePO$_4$H$_2$O

STXM measurements on CePO$_4$H$_2$O samples reacted with catechol are presented in Fig. 8. SVD analysis of a stack acquired around the C K-edge reveals that the organic material associated with CePO$_4$H$_2$O in the presence of oxygen are typical of the complex molecules produced by CeO$_2$ mediated oxidative polymerization of catechol. Organic material that is not associated with the mineral grains exhibits the lineshape of untransformed catechol. The STXM derived Ce NEXAFS spectra from the mineral did
not reveal any presence of Ce(IV). Those spectra were acquired in transmission mode, a bulk sensitive geometry. When compared with surface sensitive TEY spectra, these data indicate that CeO$_2$ formation and subsequent catechol polymerization occurs mostly at the surface of CePO$_4$/C$_1$H$_2$O grains.

3.5. Organic reaction products

Our results show that the dissolution of CePO$_4$/H$_2$O by catechol under oxic conditions at pH 5 leads to the formation of dark colored refractory material. This is true after a few days and near-complete coating of mineral particles after 10 days. By comparison, dark coloration did not appear in a stirred control catechol solution at pH 5 exposed to air until at least 500 h after preparation. In other words, direct inorganic reaction of catechol with dissolved oxygen cannot account for by coloration in the CePO$_4$/H$_2$O dissolution experiments due to variation in C speciation. The dark material was found after reacting CePO$_4$/C$_1$H$_2$O and catechol was found to be similar in structure to humic material as evidenced firstly by infrared spectroscopy (Cervini-Silva et al., 2005), and discussed in the present work. Previous reports show that strongly-sorbed humified products formed following reaction of birnessite with catechol, suggesting that such polymerized materials may be common products of mineral surface oxidation of catechol (Machjer et al., 2000). Standard reduction potential magnitude values for manganese oxides and CeO$_2$ are 1.50 and 1.61 V, respectively (Larson, 1998; Sparks, 2003).

Supernatant solution analyses confirmed the presence albeit small of oxalate and malate. Our results contrast with a previous report on the quantitative conversion of catechol to malate and oxalate after reaction with commercial CeO$_2$ at 170 °C (Duprez et al., 1996). Data discussed herein shows that surface-formed, small-sized CeO$_2$ following CePO$_4$/H$_2$O reaction induce the polymerization of organic C. Organically-coated CeO$_2$ nanoparticles have a modified surface structure (Wu et al., 2000) and can exhibit quantum size effects (Yin et al., 2002), which appear to influence mineral reactivity.

Early work (McBride, 1989a) on the transformation of catechol in the presence of birnessite at pH 5.4 reported that oxygen is not consumed during reaction. Those results were explained because Mn(IV) served as the primary
oxidant. Along these lines, given the reduction potential values for catechol and Ce(IV) and O₂, 
\[ E_{\text{Ce(IV)/Ce(III)}} = 1.61 \text{ V}, \quad E_{\text{O}_2(\text{singlet})/\text{O}_2} = 0.83 \text{ V}, \quad \text{and} \quad E_{\text{catechol} = \text{catechol}} = 0.53 \text{ V} \] (Sofen et al., 1979), it is then plausible that Ce(IV)
could also serve as primary electron acceptor towards catechol.

We monitored the dissolved oxygen concentration during the reaction between CePO₄·H₂O and catechol at 
\( p\text{H}_0 = 5 \) under oxic conditions. At \( t = 0 \), dissolved oxygen and catechol initial concentrations were \( 0.25 \pm 0.1 \) and \( 1 \pm 0.003 \) mM, respectively. After 300 h reaction time, the dissolved oxygen concentration was found to remain invariant. These results imply that the activation energy barrier for the oxidation of catechol by molecular oxygen, a diradical species, under prevailing reaction conditions (4:1, oxygen:catechol molar ratio and at room temperature), becomes limiting.

It is worth noting that a significantly higher conversion of catechol under oxic conditions in the presence of nanocrystalline CeO₂ formed on CePO₄·H₂O (The solid concentration was 1 g L⁻¹ (c) and the specific surface area (\( S_a \)) as determined by the BET method was 66 ± 1 m² g⁻¹) than in the presence of commercial CeO₂ (c = 1 g L⁻¹ and \( S_a = 19.7 \pm 1 \text{ m² g}^{-1} \)). The reaction of catechol with CePO₄·H₂O for 200 h led to the formation of CO₂ (73–85% wt C), polymeric C (8–10% wt C), and malate, oxalate, soluble catechol, and sorbed non-polymeric catechol (2–3% wt C). In contrast, the reaction between catechol and CeO₂...
under similar reaction conditions led to portioning of catechol, namely sorbed, non-polymeric (<15% wt C) and dissolved catechol (85% wt C <), with no apparent signature of catechol transformation following the mechanisms mentioned above. These results strongly suggest that (i) Ce(IV) serves as ultimate electron acceptor and (ii) Ce-surface redox properties depend on particle size.

3.6. Proposed mechanism for catechol degradation by Ce(IV)

A plausible degradation pathway for catechol by CePO₄H₂O under oxic conditions consistent with our observations involves coupled redox transformations of carbon and cerium at the mineral surface (Figs. 4–8). Oxidation of Ce³⁺ could involve formation of ternary complexes of Ce³⁺ ions, O₂, and catechol, analogous to previously described catechol–oxygen–Mn⁴⁺ complexes (Morrison and Sawyer, 1977; Magers et al., 1978; Pierpoint and Buchanan, 1981; Baes and Bloom, 1989; Sheriff, 1992; Gelasco et al., 1998). Complex formation results in a dramatic decrease in redox potential of Ce³⁺, as indicated by the potential values for Ce before and after complexation by catechol, where \( E_{\text{Ce}(IV)/\text{Ce}(II)} = +1.61 \text{ V} \) and \( E_{[\text{Ce}(cat)]^{4+}/\text{Ce}(cat)]^{3+}} = -0.448 \text{ V} \) (Sofen et al., 1979). Thus, the complexed Ce³⁺ can be rapidly oxidized by O₂.

The production of CO₂, oxalate, malate, and more complex forms of organic carbon following the reaction between CePO₄H₂O and catechol indicates that catechol undergoes oxidation and oxidative polymerization because aromatic-ring rupture (Stevenson, 1994). Charge is transferred from catechol to Ce(IV) ions, leading to semiquinone-based radicals (Stevenson, 1994) that polymerize. Finally, the oxidation of Ce³⁺ to Ce⁴⁺ is accompanied by an increase in the rate of dissolution of CePO₄H₂O as determined by the rate of accumulation of phosphate in solution (Fig. 1).

4. CONCLUSIONS

Here we report a previously unrecognized pathway of cerium biogeochemical cycling that can impact both phosphate and organic carbon cycles in terrestrial ecosystems. It is well known that phosphorous availability is impacted by changes in the redox status of soils and sediments. For instance, under reducing conditions typical of flooded rice fields or in lake sediments, minerals such as strengite [FePO₄·2H₂O], amorphous ferric phosphate [FePO₄], and trivalent manganese phosphate [MnPO₄·1.5H₂O], undergo reductive dissolution, with the release of P as a consequence (Bingham and Garber, 1960; Bingham, 1963; Moore et al., 1998; Moore, 1999; Moore and Reddy, 1999; Uusitalo and Turtola, 2003). Our results show that cerium redox transformations influence phosphorous bioavailability. In the case of CePO₄H₂O, phosphate ions are liberated into solution by cerium oxidation and the sequestration of Ce⁴⁺ in an insoluble phase (CeO₂).

In this paper, we have studied the reaction between CePO₄H₂O and catechol at pH₀ = 5 using Scanning Transmission X-Ray Microscopy. Measurements at the C K and Ce M₄,₅ edges that indicate three dominant organic species, varying in complexity and association with the inorganic phase. Untransformed catechol is loosely associated with CeO₂, whereas more complex organic molecules that exhibit lower aromaticity and stronger C=O π° resonances of carboxyl-C and carbonyl-C groups are only found in association with the grains. These results further serve as the basis to postulate that, in the presence of O₂, CeO₂ can mediate the oxidative polymerization of catechol to form higher molecular weight polymers. Also, in this paper we provide NEXAFS spectroscopic analysis to show that, in the presence of catechol, O₂ (aq) causes both the oxidation of structural Ce³⁺ and the transformation of catechol to more complex organic molecules.

The higher conversion of catechol under oxic conditions in the presence of nanocrystalline CeO₂ formed on CePO₄H₂O compared to commercial CeO₂ strongly suggest Ce(IV) serves as ultimate electron acceptor and that surface redox properties depend on particle size. Likely, energetic considerations intrinsic of changes cerium-surface redox properties such as conduction band shifting follow structural Ce oxidation, i.e., mineral transformation, affect kinetics and thermodynamics of intermolecular interactions between carbon species, mineral phases, and confined and bulk water.

In seawater, oxidation of Ce³⁺ by O₂ is mediated by microorganisms (Moffett, 1990, 1993). Thus, it has been proposed that Ce³⁺ (aq) requires microbial mediation. Data presented herein show that an alternative pathway for Ce³⁺ oxidation is feasible. Our results presented herein are explained because a pathway of biologically-induced, non-enzymatic oxidation of cerium and formation of small CeO₂ particles at room temperature. Considerations of the outcome of this work will serve as a basis for the development of a conceptual model for the use of laboratory based experimental studies with sources of organic carbon to interpret the biological fractionation of Ce in paleosols.

Our findings may be also relevant to assessment of the environmental effects of cerium compounds used in technology. CeO₂ is manufactured for a wide range of applications, including gas sensors, pigments, and as catalysts for redox reactions in automotive exhaust (Trovarelli, 1996). Nanoparticulate CeO₂ has been tested recently as a fuel additive to catalyze combustion reactions between diesel and air (Trovarelli, 1996). Catalyst cartridges bearing commercial CeO₂ have been shown to efficiently reduce the concentration of NOₓ, unburned hydrocarbons, and CO in car exhausts under real running engine conditions (Chung et al., 2003). These applications may lead to the introduction of fine grained, reactive CeO₂ into the environment, where it is capable of the low-temperature decarboxylation and polymerization of organic compounds. Consequently, our results highlight the need to study the biogeochemical and toxicological consequences of the release of CeO₂ nanoparticles into the environment.

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