The Earlier the Better: Structural Analysis and Separation of Lanthanides with Pyrroloquinoline Quinone

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Abstract: Lanthanides (Ln) are critical raw materials, however, their mining and purification have a considerable negative environmental impact and sustainable recycling and separation strategies for these elements are needed. In this study, the precipitation and solubility behavior of Ln complexes with pyrroloquinoline quinone (PQQ), the cofactor of recently discovered lanthanide (Ln) dependent methanol dehydrogenase (MDH) enzymes, is presented. In this context, the molecular structure of a biorelevant europium PQQ complex was for the first time elucidated outside a protein environment. The complex crystallizes as an inversion symmetric dimer, Eu2PQQ+, with binding of Eu in the biologically relevant pocket of PQQ. LnPQQ and Ln1Ln2PQQ complexes were characterized by using inductively coupled plasma mass spectrometry (ICP-MS), infrared (IR) spectroscopy, 151Eu Mössbauer spectroscopy, X-ray total scattering, and extended X-ray absorption fine structure (EXAFS). It is shown that a natural enzymatic cofactor is capable to achieve separation by precipitation of the notoriously similar, and thus difficult to separate, lanthanides to some extent.

Introduction

Rare earth elements (REE) include the elements 21Sc, 54Y and 57La, in addition to the 14 lanthanides (Ln) from 54Ce to 57Lu. Due to their extensive usage in modern technologies, they are also called “vitamins, or seeds of technology” and the global demand of rare earth oxides is growing steadily.[1] Unlike their name suggests, REE are not particularly rare and the occurrence of the two least abundant ones, Tm and Lu, is even higher than the one of silver.[2] Mining of those elements is, however, a challenge, due to their dispersion and low concentrations in REE containing ores. In addition, extraction methods include strong acids or bases and produce large scales of radioactive and heavy metal contaminated waste.[3] Separations of the chemically similar REE are energy-intensive and challenging.[4] However, several exciting new directions for REE separations have been presented recently. The group of Schelter used the size-sensitive ligand TriNox+ (Figure 1B), which is able to form a self-associative equilibrium out of REE mixtures and can be used for REE separation by leaching.[5a] Sun, Bünzli and co-workers used a supramolecular approach with a tris-tridentate ligand, which forms 4-nuclear cages preferentially with the smaller, late REE (Figure 1A).[6]

With a modification of the ligand, using long alkyl chains, the group was able to perform a liquid-liquid extraction of late REE, while the early ones remained in the aqueous phase.[8] Recently, also magnetic field driven REE separations have been reported.[9] Among the REE, the early lanthanides (La–Eu) are now recognized as biorelevant for methylotrophic bacteria habituating a number of different ecosystems (plant phyto-spheres, volcanic mudpots, soil and aquatic environments).[10] Those bacteria use Ln containing enzymes (methanol dehydrogenases, MDH, active site shown in Figure 2A) in their C1 metabolism. The active site of the Ln-containing enzymes includes redox cofactor PQQ (Figure 2B) that coordinates the central metal in a tridentate fashion.[11]
Remarkably, early lanthanides are taken up more quickly and preferentially by bacteria than the later ones. It was shown that *Methylorubrum extorquens* AM1 can even grow with Nd-containing hard-drive magnets as the only source of Ln, making those bacteria interesting for bioleaching or biomineralization.[14] With early Ln, bacteria grow faster and their respective MDH enzymes are more efficient in turning over methanol. Seemingly, natural systems have been tuned specifically by evolution to work best with the earlier, larger, and more abundant lanthanides. The reasons for the preference of natural systems for early lanthanides remain somewhat elusive. However, factors such as changing coordination numbers across the Ln-series, lack of efficient activation and negative impact on redox cycling of PQQ by certain Ln in the active site have been proposed.[14] PQQ is one of the few pincer ligands existing in nature and coordinates via a carboxylic acid moiety, a pyridyl nitrogen and a quinone oxygen atom.[15] Similar binding motifs have been used in ligands employed in the separation of lanthanides and actinides. 2,6-pyridine dicarboxylic acid (PDCA) and derivatives have been widely used for solvent extraction and ion chromatography.[16] The tris-tridentate ligand shown in Figure 1A also features an ONO binding pocket. Here, we show that the MDH-cofactor PQQ by itself preferentially forms complexes with early Ln and we evaluate whether this could be exploited for Ln-separation. In addition, we report the first crystal structure of PQQ with a biorelevant metal ion (Eu) outside of a MDH protein environment and without synthetic co-coordinating ligands or modified PQQ precursors.

**Results and Discussion**

**Ln1PQQ complexes: characterization**

In MDH, PQQ acts as a tridentate ligand for lanthanides, using 7'-O; 6-N and 5-O (see numbering in Figure 2B). Without the protein environment, all carboxyl groups, as well as 4-O can participate in metal coordination, often complicating analyses.[17] From aqueous solutions of the sodium salt of the cofactor (Na₃PQQ, 2, isolated from vitamin capsules) complexes with lanthanides rapidly precipitate after mixing, showing 1:1 stoichiometry, even with lanthanides added in excess (6 equiv).[17a] Single crystals from the first Eu₃PQQ complex outside the MDH protein environment were obtained for X-ray structural analysis (Figure 3). To this end, very few crystal structures of PQQ complexes have been reported[18] and none of them contained metals relevant for MDH activity (Ca, Ln). A structure of a 1:1 complex with a PQQ derivative (here the trimethylster PQQMe₃ and copper(I) were used) was reported by Kaim.[18a] Kojima reported a structure of PQQMe₃ with Ru²⁺ bearing a terpyridine coligand. A structure of PQQ (1) with Cu²⁺ and the same coligand was reported by Suzuki.[18b,c] Crystals of an Eu₃PQQ₂ complex were derived after several days from a mixture of aqueous Na₃PQQ and EuCl₃ solutions at 80 °C, which was allowed to slowly cool down to room temperature (see Supporting Information). The Eu-structure is consistent with a 1:1 stoichiometry but surprisingly reveals a dimer with head to tail coordination of PQQ. While no other co-coordinating ligands other than water were necessary to crystallize the complex, a carboxylic acid of a second PQQ molecule is needed to complete the coordination sphere. Modified PQQ derivatives, where the carboxylic acid moiety at the pyrrole ring is either blocked by alkylation to yield an ester or replaced entirely by

![Figure 1](image1.png)

**Figure 1.** A) Tris-tridentate ligand and supramolecular Ln complex reported by Sun, Bünzli et al.[15] and B) H₃TrNOx ligand and Ln TrNOx THF complex from Schelter et al.[15]

![Figure 2](image2.png)

**Figure 2.** A) Active site of a Ln-dependent MDH (PDB 6FKW). B) Tris-tridentate ligand and supramolecular Ln complex reported from Schelter et al.
for example, a methyl group, could favor the formation of mononuclear complexes that could find use as biomimetics for MDH. The ligand is fully deprotonated (PQQ$^2-$), which is not surprising as the pH of the carboxylic acids are all below the pH of the experiment. In addition to the coordination in the biologically relevant pocket (using 7'-O; 6-N and 5-O with numbering scheme in Figure 2) Eu is coordinated with 2'-O by a second PQQ molecule. The coordination sphere is completed by five water molecules (consistent with elemental. Analysis of the bulk material) yielding a final coordination number (CN) of 9. The PQQ-water adduct (C5-diol, 3), which is present to some extend in aqueous PQQ solutions, is not observed in the Eu$_2$PQQ$_2$ complex.$^{[19]}$ To gain more insight into the coordination chemistry and exclude redox processes throughout complex formation, we recorded a $^{153}$Eu Mössbauer spectrum of the precipitated bulk material. The experimental Mössbauer spectrum of Eu$_2$PQQ$_2$ at 5 K is shown in Figure 4 together with the transmission integral fit. The spectrum clearly indicates the presence of only Eu$^{3+}$ in this sample, supporting the oxidation state assigned in the crystal structure. Due to increased electron density at the core, Eu$^{3+}$ complexes commonly display negative isomer shifts around $-10$ to $-14$ mm s$^{-1}$$^{[19]}$. Here, the isomer shift of Eu$_2$PQQ$_2$ of $\delta = 0.32(2)$ mm s$^{-1}$ is well in the range for reported Eu$^{3+}$-complexes with N- and O donors such as [Eu(acac)$_3$(H$_2$O)$_3$] ($\delta = 0.36$ mm s$^{-1}$, at 80 K), [Eu(NO)$_3$(phen)$_3$] ($\delta = 0.41$ mm s$^{-1}$, at 80 K) or H[Eu(dicta):4H$_2$O ($\delta = 0.30$ mm s$^{-1}$, at rt) and Ba[Eu(dtpa)]:7.5H$_2$O ($\delta = 0.38$ mm s$^{-1}$, at rt).$^{[20]}$ The fitting of the Eu$_2$PQQ$_2$ data further reveals an electric quadrupole splitting of $\Delta E_Q = 1.6(4)$ mm s$^{-1}$ and an experimental line width of $\Gamma = 2.5(1)$ mm s$^{-1}$. The small experimental line width indicates one single europium species rather than multiple Eu$^{3+}$ ions in crystallographically distinct sites.$^{[19h,20a]}$ Usually for Eu$^{3+}$ complexes (and other Eu$^{3+}$ solids), the range of observed isomer shifts is only within 0.8 mm s$^{-1}$ and largely insensitive to the nature and number of coordinating ligands.$^{[20a]}$ However, the Mössbauer spectroscopic data supports overall the presence of only one Eu$^{3+}$ species in the bulk material.

To analyze whether the structure shown in Figure 3 is representative for the bulk material that precipitated and to compare coordination numbers (CN) and metal-ligand distances of different LnPQQ complexes (Ln = La, Eu, Tb, Lu), we turned to EXAFS spectroscopy and pair distribution functions analysis (PDF). The results are shown in Table S2, Figure 5 and in the Supporting Information. Among four evaluated monometallic species, only EuPQQ showed good crystallinity with relatively sharp diffraction lines in the powder X-ray diffraction (Figure 5A). The diffractogram characteristic for bulk material can be reconstructed up to experimental $Q_{max} < 2.3$ Å$^{-1}$. Nevertheless, interatomic correlations for LuPQQ, TbPQQ and LaPQQ (Figure 5B) can be detected only up to $r = 5$–8 Å, which is typical for amorphous materials without long-range order and without detectable M···M correlations at approximately 12–14 Å obtained for Eu$_2$PQQ$_2$. Qualitatively, the short-range structure characteristic for TbPQQ and LuPQQ is similar. Average Ln–L distances (Table S2) are similar and the coordination number is comparable for all three species. The LaPQQ local structure seems to differ slightly from EuPQQ with visibly longer La–L distances.
Analysis of the EXAFS data showed that from early to late Ln, both the CN as well as average Ln–L distances are reduced (Table S2 and Figure 6). The determination of the coordination number was achieved by using the well-established data reduction procedures presented by Ravel et al. A first shell fit was performed between the measured LnPQQ EXAFS and the model based on Eu₄POQ single-crystal data. The first shell fit entails the scattering paths from all neighboring atoms presented in Table S2, obtained from the used model. This comprises one Ln–O1 at 2.36 Å, two Ln–O9 at 2.423 Å, three Ln–O1 at 2.45 Å, one Ln–O12 at 2.48 Å, one Ln–O3 at 2.57 Å and one Ln–N1 at 2.65 Å. This results in a total coordination number of 9. In the case of LaPQQ and Eu₄PQQ this fitted well, but in the case of TbPQQ and LuP QQ a total coordination number of seven yielded a better fit. The total coordination number results in the first peak observed experimentally (Figure 6), entailing all scattering paths, supported by the model. Furthermore, the position of this peak delivers information about the Ln–L distance. Measured EXAFS supported by the model show different average ligand distances as in Table S2. La–L is at 2.64 Å, Eu–L at 2.48 Å, Tb–L at 2.44 Å and Lu–L at 2.35 Å.

The coordination sphere of Eu (CN = 9) in the 1:1 Eu₄PQQ and LaPQQ complex would therefore consist of five equiv of water and four times coordinated PQQ. For the later Ln, Tb and Lu, the CN is reduced concurrent to the number of coordinated water molecules. This agrees with the first shell fit by EXAFS, where a CN of 7 is obtained (for further details, see Methods in Supporting Information). The decrease of the preferred CN throughout the lanthanide series is a common feature in lanthanide coordination chemistry. Analytical data of a previously reported LaPQQ sample was in best agreement with five equiv of water. A correlation is further recognized between the determined CN with XAS and the calculated water equivalents based on elemental analysis on the one hand and the IR intensity of the large O–H stretching feature between 3600-2500 cm⁻¹ (Figure S1).

By comparing IR spectra of the 1:1 LnPQQ complexes (Ln = La, Eu, Tb, Lu) with Na₄PQQ·H₂O large similarities are revealed, which indicate participation of at least two of the three carboxyl groups in Ln coordination (Figure S1) as also found in the Eu₄POQ structure. In comparison with POQ (1, > 20 main peaks) the complexes exhibited IR spectra with only eight main features resolved: 3630–2500, 1735, 1580, 1507, 1351, 1241, 1195 and 1149 cm⁻¹. Especially the seven quinone and carboxyl related vibrations of POQ merge to a broad, poorly resolved area between 1735–1580 cm⁻¹. Further, the IR spectra of the LnPQQ complexes reveal more similarities to Na₄POQ than to free POQ (Figure S1). More importantly, the IR spectra of bulk material and single crystals of the Eu₄POQ dimers show almost identical IR spectra (Figure S2), further confirming that the obtained crystal structure is representative for the bulk material.

**Ln1Ln2PQQ complexes: structure and lanthanide preference**

The property of aqueous solutions of Na₄PQQ to instantly precipitate one equivalent of lanthanide prompted us to study whether there would be a preferred Ln when mixtures of Ln were offered. Considering the demand of new separation techniques for REE and the preference of bacteria for certain lanthanides, we were curious about the ability of POQ to separate mixtures of Ln. Importantly, POQ does not have to be made synthetically but can be obtained in kg amounts by fermentation without the use of any organic solvents and can further be fully recovered and separated from Ln by addition of concentrated HCl. This will destroy the complex and lead to precipitation of fully protonated POQ (1) which is extremely stable against acids. In fact, it needed a mixture of boiling concentrated nitric and hydrochloric acid to fully dissolve (and possibly decompose) 1 in our studies. Stock solutions of Na₄PQQ in water were mixed with aqueous solutions of the lanthanide chlorides, each consisting of three equiv of lanthanum chloride and three equiv of a second lanthanide chloride. All samples showed similar IR spectra compared to the monometallic LnPQQ complexes (Figure S3). And like the LnPQQ species, small differences appeared in the height of the large IR-absorption band between 3630–2500 cm⁻¹, stemming from different equiv of coordinated (or cocrystallized) water (Figure S3, Table S3). Analysis of the received precipitated material still revealed a 1:1 PQQ/metal stoichiometry by elemental analysis and ICP-MS, but the metal content was now distributed between lanthanum and the additional Ln (see Supporting Information). The experiments were repeated three times (experimental replicates) and the distribution between the two added metals in the 1:1 LnPQQ complexes is given in Figure 7 in percent.

Remarkably, when the additional metal is an early lanthanide (Ce–Eu) its uptake into the complex is preferred over lantha-
num to up to 57% for Ce, but slowly decreases over the series to 53% for Eu. Preference for the second lanthanide then further decreases, falling below La, starting from Gd (46%) to 26–28% for the last three lanthanides (Tm–Lu). In general, the slightly different properties of Ln caused by lanthanide contraction usually cause increasing complex stabilities from La to Lu,[22b] concurrent with higher affinities to ligands such as EDTA.[4] Exceptions with rigid macrocycles such as functionalized aza-crown ethers, where the larger Ln form more stable complexes are rare but have been reported.[25] It is interesting that the “break” between favoring early Ln (Ce–Eu) over La versus favoring La over late Ln (Gd–Lu) occurs at Gd. Notably, in activity measurements of PQQ- and Ln-dependent MDH isolated from *M. fumaroliicum* SolV, it was also shown that activity was not stimulated anymore with the later lanthanides starting with Gd.[14]

Due to their different Lewis acidities, the lanthanide stock solutions exhibited varying pH values (pH 4.8–6.1, MilliQ water pH 5.9). Hence, to rule out an effect of varying pH, an additional experiment with fixed pH at 3.5 was performed. In the Ln series at 25 °C, ICP-MS of the precipitated solids revealed a similar pattern along the series as without controlled pH, although the percentage of lanthanum was overall slightly increased when an equimolar mixture of lanthanum and the following series at 25 °C was tested again in this series as control, as well as against each other, as well as the Gd/Tb pair. The La/Ce pair was tested against each other, as well as the Gd/Tb pair. The La/Ce pair was tested against each other, as well as the Gd/Tb pair. The La/Ce pair as separation factor SF was determined. Both Eu in the Eu/Gd couple and Gd in the Gd/Tb couple show a ≈ 10% higher incorporation over the following Ln in the series. With larger size differences between the two REE, the difference in incorporation in the solid phase increases. With Eu and Y, the two main elements of YOX, the precipitated EuYPQQ solid contains only 19% of yttrium. With equimolar amounts of Nd and Dy, only 29% of the smaller Dy gets coordinated by, and precipitates with PQQ. For the latter pair a separation factor SF of 2.4 was determined.

To investigate whether the separation is possibly based on structural preference, preorganization and size of the Ln/REE, the structures of the Ln1Ln2PQQ bulk materials were analyzed with the same techniques used for the monometallic LnPQQ complexes discussed above. The bimetallic EuLaPQQ complex shows high crystallinity with a powder X-ray diffraction pattern similar to monometallic EuPQQ. The diffraction pattern of Eu-LaPQQ can also be completely indexed using the cell parameters and space group obtained for the complex shown in Figure 2. TbLaPQQ and LuLaPQQ are amorphous (Figure S5A). Their pair distribution functions do not show correlations above r = 8 Å (Figure 8). The PDF of TbLaPQQ is similar to TbPQQ but does not fit with LaPQQ, while the PDF for LuLaPQQ seems to be a superposition of profiles characteristic for LaPQQ and LuPQQ, indicating that LuLaPQQ might be a mixture of two species or might have completely different structure in comparison with LaPQQ and LuPQQ with possible ordering of metals in structure or islands occupied by different metals.
Conclusions

In this study, we obtained the first crystal structure of PQQ with a lanthanide ion (Eu$^{3+}$) and without the need of any co-ligands. Different spectroscopic methods were used for extensive characterization of this and other LnPQQ complexes. Here, we also find that complexes with early lanthanides are preferentially formed with PQQ and rapidly precipitate out of aqueous solutions. It is very likely that the size of the Ln plays a role in the observed separation. While separation of Ln with PQQ is not competitive with existing processes at present, it is nevertheless noteworthy that a natural enzymatic cofactor is capable to achieve separation of the notoriously similar and thus difficult to separate Ln to some extent. Remarkably, Ln-utilizing bacteria also show a preference for early lanthanides (uptake and growth), and isolated methanol dehydrogenases are more active with earlier Ln. It has been reasoned that the size and coordination of the Ln in the active site might be responsible for this, corroborated both by molecular dynamics simulations combined with fragment molecular orbital as well as with DFT calculations.\textsuperscript{14,28} We hope that the thorough structural analysis described here will also help to characterize the exciting, recently discovered natural lanthanide enzymes to shed light into the many open questions in this emerging field of lanthanide-dependent bacterial metabolism.

Experimental Section

Crystallographic data: Deposition number 1991651 contains the supplementary crystallographic data for this paper. These data are provided free of charge by the joint Cambridge Crystallographic Data Centre and Fachinformationszentrum Karlsruhe Access Structures service.

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Conflict of interest

The authors declare no conflict of interest.

Keywords: coordination chemistry · lanthanides · PQQ · rare earth elements · separations

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