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Structures of hydrated and hydrolyzed thorium(IV) determined by EXAFS and LAXS

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The hydrated thorium(IV) ion is nine-coordinate in strongly acidic aqueous solution with a mean Th-O bond distance of 2.45(1) Å as determined by extended X-ray absorption fine structure (EXAFS) and large angle X-ray scattering (LAXS) [1]. The presence of a second hydration sphere was indicated by LAXS with cca 18 water molecules at 4.66(2) Å. Structural studies in aqueous solution of the hydrolysis products of thorium(IV) identi $fied three different types of hydrolysis species: a \mu < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > (OH) < sub > 2 < / sub > (H < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > (OH) < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo dimer, [Th < sub > 2 < / sub > O-hydroxo d$ $a\ \mu < sub > 2 < / sub > O - hydroxo\ tetramer, [Th < sub > 4 < / sub > (OH) < sub > 8 < / sub > (H < sub > 2 < / sub > O) < sub > 16 < / sub >] < sup > 8 + < / sup > (H < sub > 2 < / sub > O) < sub > 16 < / su$ $and\ a\ \mu₃O-oxo\ hexamer, [Th₆O-sub>8</sub>(H₂O)_n]⁸⁺.$ Strongly acidic solutions are required to stabilize the hydrated thorium(IV) ion. With increasing concentration and pH thorium(IV) hydrolyzes, first, to a dimer with double hydroxo bridges, then to a tetramer with the thorium(IV) ions forming a diamond (represented by two dimers merged together) [1]. Finally, at pH value close to precipitation of thorium(IV) oxide/hydroxide, a hexamer is formed with the thorium atoms forming a rhomboctahedron. Potentiometric studies suggested a hexamer composition [Th₆(OH)_x(H₂O)_nn</sub> x</sup>, x = 14 or 15[2]. However, the rhomboctahedral structure prevents for sterical reasons such a large number of hydroxide groups to be bound. Instead 7-8 oxo groups are bound to thorium ions as from proton activity point of view 16 hydroxo groups are equivalent to 8 oxo groups, making it impossible to distinguish between e.g. Th₆(OH)₁₆⁸⁺ and Th₆O₈⁸⁺ complexes. A compilation of reported solid state structures of actinoid(IV) compounds with oxygen donor ligands show a strong correlation between the An-O bond distance and the coordination number [1]. EXAFS studies of uranium(IV), neptunium(IV) and plutonium(IV) ions in aqueous solution have reported accurate An-O bond distance, but the coordination numbers were less accurate and often overestimated. The hydrated tetravalent actinoid ions in aqueous solution all seem to be nine-coordinated [1].

References

- 1. Torapava, N., Persson, I., Eriksson, L., Lundberg, D.: Inorg. Chem. 48, 11712 (2009)
- 2. Rand, M., Fuger, J., Grenthe, I., Neck, V., Rai, D.: Chemical Thermodynamics of Thorium, Eds. Mompean, F. J., Perrone, J., Illemassène, M., OECD Nuclear Agency Agency, Data Bank, F-92130 Issy-les-Moulineaux, France, North Holland Elsevier Science Publishers B. V., Amsterdam, The Netherlands 2008, and references therein.

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