OPEN ACCESS Crystals ISSN 2073-4352 www.mdpi.com/journal/crystals

Article

The First Lanthanide Telluride-Bromide: La₃Te₄Br, a Valence Compound

Markus Larres¹, Anja-Verena Mudring² and Gerd Meyer^{1,*}

- ¹ Department für Chemie, Universität zu Köln, Greinstraße 6, D-50939 Köln, Germany; E-Mail: larresm@uni-koeln.de
- ² Fakultät für Chemie, Ruhr-Universität Bochum, D-44780 Bochum, Germany;
 E-Mail: anja.mudring@rub.de
- * Author to whom correspondence should be addressed; E-Mail: gerd.meyer@uni-koeln.de; Tel.: +49-221-470-3262; Fax: +49-221-470-5083.

Received: 21 February 2011; in revised form: 10 March 2011 / Accepted: 18 March 2011 / Published: 21 March 2011

Abstract: The first ternary lanthanide telluride-bromide La₃Te₄Br was obtained from a mixture of LaTe and LaBr₃ (in the presence of iridium) in a sealed tantalum container at elevated temperatures. The crystal structure (orthorhombic, *Pnma*, *a* = 1634.3(2), *b* = 435.0(1), *c* = 1426.6(2) pm, Z = 4, R₁ (I₀ > 2 σ (I₀)) = 0.0349) is built from dicapped trigonal prisms of tellurium and bromine atoms surrounding lanthanum in two different ways. The dicapped trigonal prisms are connected via common edges to a three-dimensional structure, in the same fashion as is known for the binary U₃Te₅. La₃Te₄Br is a valence compound according to (La³⁺)₃(Te²⁻)₄(Br⁻) and one out of three lanthanide telluride-halides known to date.

Keywords: Lanthanum; Telluride-bromide; Synthesis; Crystal structure

1. Introduction

There is a remarkable difference in our knowledge of chalcogenide-halides RChX of the rare-earth elements (R) when the chalcogen (Ch) part is considered alone; while the oxide-halides ROX, the most prolific compound type, are all known and were structurally characterized at least by powder X-ray diffraction, in many cases by single crystal X-ray structure determination, see for example ref. [1],

telluride-halides are almost unknown. It appears that only two compounds in the R/Te/I system have been well characterized, virtually none in the other R/Te/X systems. The two R/Te/I compounds, $Er_{17.3}Te_{24}I_8$ [2] and La_2TeI_2 [3], also mark the two principal alternatives in telluride-halide chemistry. The first, a three-dimensional nonstoichiometric compound, may be derived from the NaCl-type of structure with polytelluride fragments, with tellurium in the oxidation states -2 and -1. The latter, La_2TeI_2 , would be counted as a metal-rich layer compound with condensed octahedral clusters; it has the same structure as $3R-Lu_2CCl_2$ [4], better represented as { CLu_2 } Cl_2 = { $TeLa_2$ } I_2 [5]. Octahedral lanthanum clusters with endohedral tellurium atoms, { $TeLa_{6/3}$ }, share common edges to double layers which are sheathed by closest packed iodide layers above and below.

Rare-earth element clusters with endohedral atoms of both main-group and transition elements are a steadily growing class of compounds which must be seen in connection with the cluster chemistry of the early transition elements, especially groups 4 through 6 [5-20]. The configuration crossover $4f^{n}5d^{0} \leftrightarrow 4f^{n-1}5d^{1}$ of some of the lanthanides allows for bonding interactions between, predominantly, the endohedral atom Z and the cluster atoms R [5,16-19].

There is also a growing number of metal-rich lanthanide tellurides, see especially [18], of which Sc_2Te [20] was one of the first binary examples. The ternary compound $Sc_{14}Os_3Te_8$ [21] links this chemistry structurally to the reduced rare-earth element chemistry just mentioned. $Sc_{14}Os_3Te_8$, or better $\{Os_3Sc_{12}\}Te_8Sc_{2-x}$, has remarkable structural similarities with $\{Os_3Sc_{12}\}Br_{16}Sc$ [22]. Both compounds contain the same extended cluster chain of square antiprisms and cubes of scandium atoms (in a 2:1 ratio) with endohedral eight-coordinate osmium atoms! The chains are surrounded by eight telluride (-2) and sixteen bromide (-1) ions, respectively, and additional scandium atoms reside in octahedral interstices between the hexagonally close-packed chains.

One goal of our present research in this area is to combine these two chemistries and create metalrich condensed-cluster telluride-halides. Consequently, first explorations in the systems Z/R/Te/X were designed for compounds like {Ir₃R₁₂}Br₈Te₄. Although we have not yet accomplished our goal, we have, in the Ir/La/Te/Br system, obtained the first lanthanide telluride bromide, La₃Te₄Br.

2. Results and Discussion

The ternary compound La₃Te₄Br is obtained from mixtures of LaTe, LaBr₃ and Ir at elevated temperatures in a sealed tantalum reaction container. The crystal structure of La₃Te₄Br (orthorhombic, *Pnma*, a = 1634.3(2), b = 435.0(1), c = 1426.6(2) pm, Z = 4) is best regarded as a valence compound, according to $(La^{3+})_3(Te^{2-})_4(Br)$. All Te—Te distances are at or beyond 377 pm which attests for single non-bonded Te²⁻. For comparison, all bonding distances in LaTe₂—which might, roughly, be understood as a metallic ditelluride, $(La^{3+})(e^{-})(Te_2^{2-})$ [23-25]—are between 299 and 341 pm.

As all atoms occupy the Wyckoff site (4c) in the orthorhombic space group *Pnma*, there are as many crystallographically independent atoms in La₃Te₄Br as the formula claims, with four formula units in the unit cell. All three lanthanum atoms exhibit coordination number eight, with dicapped trigonal prisms as the corresponding polyhedra. La1 and La2 are surrounded by seven tellurium atoms with one tellurium and one bromine atom as the caps of two rectangular faces, and La3 is surrounded by six tellurium and two bromine atoms with the bromine atoms forming one edge of the trigonal prism. The three prisms are connected via two common edges (Te2—Te4 and Te3—Br) to a trimeric building unit

of the composition La₃Te₁₄Br₄. The *b* axis as the short axis, 435.0(1) pm, is identical with the height of the trigonal prisms. Thus, there are Te1—Te1, Te2—Te2, Te3—Te3, Te4—Te4, and Br—Br distances all of this length. Te—Te distances in the triangles and to the caps are usually shorter, starting at 377 pm and going up to 461 pm, see legend to Figure 1. The Te—Te distance in NaCl-type NaTe is 455 pm [23]. There are two bromine functions (terminal and capping) in La₃Te₄Br, which are reflected in the La—Br distances of 304.60(9) (prism edge) and 310.78(14)/312.74(14) pm (caps). These distances compare quite well with the corresponding distances in LaBr₃, 310 (6x) and 316 pm (3x) for the tricapped trigonal prism of its UCl₃-type structure [26]. La—La distances within the La₃Te₁₄Br₄ building block are 476.1(1) pm (La1—La2), 456.5(1) pm (La1—La3), and 462.4(1) pm (La2—La3).

Figure 1. The building block $La_3Te_{14}Br_4$ in the crystal structure of La_3Te_4Br : the three crystallograhically independent La atoms (La1: red; La2: yellow; La3: blue) and their surrounding by Te and Br atoms. Interatomic distances in pm and angles in degrees (with estimated standard deviations in parentheses): La1-Br1 = 310.78(14), $La1-Te2^{(i)} = 329.26(8)$, La1-Te2 = 329.26(8), $La1-Te4^{(ii)} = 329.55(8)$, $La1-Te4^{(iii)} = 329.55(8)$, $La1-Te4^{(iii)} = 329.55(8)$, La1-Te4 = 333.37(11), $La1-Te3^{(i)} = 337.46(8)$, La1-Te3 = 337.46(8), $La2-Br1^{(ii)} = 312.74(14)$, $La2-Te3^{(iii)} = 327.98(8)$, $La2-Te3^{(v)} = 327.98(8)$, La2-Te2 = 329.77(11), La2-Te4 = 330.64(8), $La2-Te4^{(vi)} = 330.64(8)$, La2-Te1 = 333.88(8), $La2-Te1^{(i)} = 333.88(8)$, $La3-Te1^{(vii)} = 325.38(8)$, $La3-Te2^{(viii)} = 328.20(8)$, $La3-Te2^{(x)} = 328.20(8)$, La3-Te3 = 332.88(11), $La3-Te1^{(xii)} = 337.01(11)$. Operators for generating equivalent atoms: (i) x, y+1, z; (ii) -x, -y, -z+1; (iii) -x+\frac{1}{2}, -y, z+ $\frac{1}{2}$; (x) -x+ $\frac{1}{2}$, -y-1, z+ $\frac{1}{2}$; (xi) -x, -y-1, -z+1.



The building blocks $La_3Te_{14}Br_4$ as exhibited in Figure 1 are connected such that each of the tellurium atoms have five lanthanum neighbors and each of the bromine atoms has four direct lanthanum neighbors, such that the connectivities might be written as follows, $La_3(Te_{7/5})_2Te_{6/5}Br_{4/4} = La_3Te_{20/5}Br_{4/4} = La_3Te_4Br$. This is achieved in a way that Figure 2 exhibits in two perspective projections, down [010] and [100], respectively.

Figure 2. Two views of the connections of the $La_3Te_{14}Br_4$ building blocks to the threedimensional structure of La_3Te_4Br ; projections down [010] (top) and [100] (bottom).



The crystal structure of La₃Te₄Br is a derivative of the structure of U₃Te₅ [27]. The latter has a remarkably smaller unit cell, owing to the mixed-valence nature of $U_3Te_5 = (U^{3+})_2(U^{4+})(Te^{2-})_5$, with both U^{3+} and especially U^{4+} considerably smaller than La³⁺.

Meanwhile, we have obtained a number of RTeI type telluride-iodides, for example PrTeI which belongs to the PbFCl-type of structure [28].

3. Experimental Section

All reactions and handling were carried out under a dry nitrogen atmosphere using dry box equipment (MBraun, Garching, Germany). LaBr₃ was prepared following the ammonium bromide route and purified as described previously [29-32]. LaTe was synthesized from a 1:1 molar ratio of La powder (Chempur, Karlsruhe, Germany, 99.9%) and Te granules (Sigma-Aldrich, München, Germany, 99.99%). The present single crystals were obtained from a reaction of LaTe (150 mg, 0.56 mmol), LaBr₃ (150 mg, 0.40 mmol) and Ir (90 mg, 0.47 mmol). The starting materials were filled into He-arc

welded tantalum ampoules jacketed with an evacuated silica tube to prevent oxidation at high temperatures. The following temperature program was applied in a tubular furnace: heated to 1,323 K with 50 K/h, kept at that temperature for 24 hours, then cooled slowly to 1,223 K (2 K/h) and annealed for 248 hours, then cooled to 973 K with 5 K/h, kept there for 1 hour, and cooled to room temperature over 8 hours. The tantalum tube was transferred to a dry-box and the contents inspected with the aid of a microscope.

La₃Te₄Br forms well-faceted, polygonal black crystals. These were selected under a microscope and sealed in thin-walled glass capillaries. After their quality had been checked by Laue diffraction patterns, the single crystals were transferred to a single-crystal X-ray diffractometer (Stoe Image Plate Diffraction System, IPDS I) to collect a complete intensity data set at ambient temperature. Structure solution and refinement was performed with the programs SHELXS-97 (direct methods) [33] and SHELXL-97 [34], scattering factors were from International Tables for X-ray Crystallography [35]. Data corrections were carried out for Lorentz and polarization factors and absorption (numerical with the aid of the programs X-RED [36] and X-SHAPE [37]). Further details of the crystal structure determination may be obtained from the Fachinformationszentrum Karlsruhe, D-76344 Eggenstein-Leopoldshafen, Germany (fax: (+49)7247-808-666; e-mail: crysdata@fizkarlsruhe.de), on quoting the depository number ICSD-380475, the authors and the journal citation.

Crystal data for La₃Te₄Br (1007.04 g mol⁻¹); diffractometer IPDS-I, Stoe, Darmstadt; Mo-K_{α} (graphite monochromator, $\lambda = 71.073$ pm); T = 293(2) K; $2\theta_{max} = 56.3^{\circ}$; 100 images, $0^{\circ} \le \phi \le 200^{\circ}$; $\Delta \phi = 2^{\circ}$; indices: $-21 \le h \le 21$, $-5 \le k \le 5$, $-18 \le l \le 18$; transmission (min, max) = 0.0307, 0.0929; $\rho_{calc} = 6.596$ g cm⁻³; 9390 reflection intensities measured of which 1318 were symmetrically independent, R_{int} = 0.0571, F(000) = 1656, $\mu = 27.547$ mm⁻¹. Orthorhombic, Pnma (no. 62), a = 1634.3(2), b = 435.0(1), c = 1426.6(2) pm, V = 1014.1(2) 10⁶ pm³, Z = 4. R values: R₁/wR₂ for 1120 reflections with [I₀ > 2 σ (I₀)]: 0.0349/0.0920 and for all data: 0.0426/0.0965; S_{all} = 1.072.

4. Conclusions

The first lanthanide telluride bromide, La_3Te_4Br , was obtained from LaTe and LaBr₃ (in the presence of iridium metal which has no effect on the formation of La_3Te_4Br) at elevated temperatures. The black crystals crystallize with a variant of the U_3Te_5 type of structure with dicapped trigonal prisms (2 × LaTe₇Br and LaTe₆Br₂, respectively) connected via common faces and edges to a three-dimensional structure. La₃Te₄Br is best considered as a valence compound according to (La³⁺)₃(Te²⁻)₄(Br⁻).

Acknowledgements

This work was supported by the Deutsche Forschungsgemeinschaft, Bonn, within the framework of the Sonderforschungsbereich 608 (Complex transition metal compounds with spin and charge degrees of freedom and disorder).

References

1. Zimmermann, S.; Meyer, G. A Missing Rare-Earth Oxide Halide Structure Now Observed For Scandium Oxide Iodide, ScOI. Z. Anorg. Allg. Chem. 2008, 634, 2217-2220.

- 2. Stöwe, K. Nonstoichiometry in Er–Te Binary and Er–Te–I Ternary Systems. *J. Solid State Chem.* **1998**, *139*, 57-65.
- 3. Ryazanov, M.; Simon, A.; Mattausch, H.J. La₂TeI₂: A New Layered Telluride Iodide with Unusual Electrical Properties. *Inorg. Chem.* **2006**, *45*, 10728-10733.
- 4. Schleid, T.h.; Meyer, G. Synthesis and Crystal Structures of Hydrogen and Carbon Stabilized Lutetium Monochloride, LuClH_x and Lu₂Cl₂C. *Z. Anorg. Allg. Chem.* **1987**, *552*, 90-96.
- 5. Meyer, G. Cluster Complexes as *anti*-Werner Complexes. Z. Anorg. Allg. Chem. 2008, 634, 2729-2736.
- 6. Corbett, J.D. Reduced Halides of the Rare Earth Elements. *Rev. Chim. Minérale* **1973**, *10*, 239-257.
- 7. Corbett, J.D. Extended Metal-Metal Bonding in Halides of the Early Transition Metals. *Acc. Chem. Res.* **1981**, *14*, 239-246.
- 8. Meyer, G. Reduced Halides of the Rare-Earth Elements. *Chem. Rev.* **1988**, 88, 93-107.
- 9. Simon, A. Clusters of Valence Electron Poor Metals—Structure, Bonding, and Properties. *Angew. Chem. Int. Ed. Engl.* **1988**, *27*, 159-183.
- Simon, A.; Mattausch, H.j.; Miller, G.J.; Bauhofer, W.; Kremer, R.K. Metal-Rich Halides. Structures, Bonding and Properties. In *Handbook on the Physics and Chemistry of Rare Earths*; Gschneidner, K.A., Eyring, L., Eds.; Elsevier Science Publishers B.V.: Amsterdam, The Netherlands, 1991; Volume 15, pp. 191-285.
- 11. Meyer, G.; Meyer, H.-J. Unusual Valences in Rare-Earth Halides. Chem. Mater. 1992, 4, 1157-1168.
- 12. Corbett, J.D. Diverse solid-state clusters with strong metal-metal bonding. In praise of synthesis. *J. Chem. Soc. Dalton Trans.* **1996**, 575-587.
- 13. Corbett, J.D. Exploratory Synthesis in the Solid State. Endless Wonders. *Inorg. Chem.* **2000**, *39*, 5178-5191.
- Meyer, G.; Wickleder, M.S. Simple and Complex Halides. In *Handbook on the Physics and Chemistry of Rare Earths*; Gschneidner, K.A., Eyring, L., Eds.; Elsevier Science B.V.: Amsterdam, The Netherlands, 2000; Volume 28, pp. 53-129.
- 15. Corbett, J.D. Exploratory synthesis of reduced rare-earth metal halides, chalcogenides, intermetallics: New compounds, structures, and properties. *J. Alloys Compd.* **2006**, *418*, 1-20.
- 16. Simon, A.; Mattausch, H.j.; Ryazanov, M.; Kremer, R.K. Lanthanides as d Metals. Z. Anorg. Allg. Chem. 2006, 632, 919-929.
- 17. Meyer, G. The Reduction of Rare-Earth Metal Halides with Unlike Metals Wöhler's "Metallothermic Reduction". Z. Anorg. Allg. Chem. 2007, 633, 2537-2552.
- 18. Corbett, J.D. Exploratory Synthesis: The Fascinating and Diverse Chemistry of Polar Intermetallic Phases. *Inorg. Chem.* **2010**, *49*, 13-28.
- 19. Simon, A. Metal clusters inside out. Phil. Trans. R. Soc. A 2010, 368, 1285-1299.
- 20. Maggard, P.A.; Corbett, J.D. Sc₂Te: A Novel Example of Condensed Metal Polyhedra in a Metal-Rich but Relatively Electron-Poor Compound. *Angew. Chem. Int. Ed. Engl.* **1997**, *36*, 1974-1976.
- Chen, L.; Corbett, J.D. Remarkable Metal-Rich Ternary Chalcogenides Sc₁₄M₃Te₈ (M = Ru, Os). J. Am. Chem. Soc. 2003, 125, 1170-1171.

- Zimmermann, S.; Brühmann, M.; Casper, F.; Heyer, O.; Lorenz, T.; Felser, C.; Mudring, A.-V.; Meyer, G. Eight-Coordinate Endohedral Rhenium, Osmium and Iridium Atoms in Rare-Earth Halide Cluster Complexes. *Eur. J. Inorg. Chem.* **2010**, 2613-2619.
- 23. Wang, R.; Steinfink, H.; Bradley, W.F. The Crystal Structure of Lanthanide Telluride and Tellurium-Deficient Neodymium Telluride. *Inorg. Chem.* **1966**, *5*, 142-145
- 24. Stöwe, K. Crystal Structure and Electronic Band Structure of LaTe₂. J. Solid State Chem. 2000, 149, 155-166.
- 25. Ramsey, T.H.; Steinfink, H.; Weiss, E.J. The Phase Equilibria and Crystal Chemistry of the Rare Earth-Group VI Systems. IV. Lanthanum-Tellurium. *Inorg. Chem.* **1965**, *4*, 1154-1157.
- 26. Krämer, K.; Schleid, T.h.; Schulze, M.; Urland, W.; Meyer, G. Three Bromides of Lanthanum: LaBr₂, La₂Br₅, and LaBr₃. *Z. Anorg. Allg. Chem.* **1989**, *575*, 61-70.
- 27. Tougait, O.; Potel, M.; Noel, H. Crystal Structure and Magnetic Properties of the Binary Triuranium Pentatelluride U₃Te₅. J. Solid State Chem. **1998**, 139, 356-361.
- 28. Larres, M.; Meyer, G. Universität zu Köln, Köln, Germany. Unpublished results, 2011.
- 29. Meyer, G.; Ax, P. An Analysis of the Ammonium Chloride Route to Anhydrous Rare-Earth Metal Chlorides. *Mat. Res. Bull.* **1982**, *17*, 1447-1455.
- 30. Meyer, G.; Dötsch, S.; Staffel, T. The Ammonium-Bromide Route to Anhydrous Rare Earth Bromides, MBr₃. J. Less-Common Met. **1987**, 127, 155-160.
- 31. Meyer, G. The Ammonium Chloride Route to Anhydrous Rare Earth Chlorides The Example of YCl₃. *Inorg. Synth.* **1989**, *25*, 146-150.
- 32. Meyer, G. Binary Lanthanide(III) Halides, MX₃ (X = Cl,Br,I). In *Synthesis of Lanthanide and Actinide Compounds*; Meyer, G., Morss, L.R., Eds.; Kluwer Acad. Publ.: Dordrecht, The Netherlands, 1991; pp. 135-144.
- 33. Sheldrick, G.M. *SHELXS-97, Program for Structure Analysis*; University of Göttingen: Göttingen, Germany, 1998.
- 34. Sheldrick, G.M. *SHELXL-93, Program for Crystal Structure Refinement*; University of Göttingen: Göttingen, Germany, 1993.
- 35. Wilson, A.J.C. *International Tables for Crystallagraphy*; Kluwer Acad. Publ.: Dordrecht, The Netherlands, 1992; Volume C.
- 36. Stoe. *X-RED 1.22, Stoe Data Reduction Program (C)*; Stoe & Cie GmbH: Darmstadt, Germany, 2001.
- 37. Stoe. *X-Shape 1.06, Crystal Optimisation for Numerical Absorption Correction (C)*; Stoe & Cie GmbH: Darmstadt, Germany, 1999.

© 2011 by the authors; licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution license (http://creativecommons.org/licenses/by/3.0/).