Stabilizing the Homopolycation (I$_4$)$_2$$^{2+}$ with a Hexasulfate in (I$_4$)[S$_6$O$_{19}$] and a Borosulfate in (I$_4$)[B(S$_2$O$_7$)$_2$]$_2$

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The formation of red and green coloured solutions when concentrated sulfuric acid reacts with tellurium and selenium, respectively, is well known and a typical show experiment in freshman courses. In fact, these reactions can be traced back to the early days of chemistry and Klaproth was the first to report the tellurium reaction.\[3,4\] Today we know that the observed colour is caused by the formation of polycations, in this case [Te$_3$]$_2$$^2+$ and [Se$_3$]$_2$$^2+$, respectively.\[5,6\] Moreover, quite a number of further chalcogen polycations have been reported up to date, significantly inspired by the work of Gillespie et al. in the early 1970s.\[5,6\] Interestingly, most of these cations are not stabilized as oxoanionic salts, although their discovery originates from reactions in sulfuric acid. Usually, the polycations are crystallized as fluoro-antimonates or -arsenates, and [Se$_3$][HSO$_4$], obtained from Se and oleum, is one of the few exceptions.\[7\] The early discovery of chalcogen cations has certainly triggered investigations aiming at the preparation of similar cations, including the heavier halogens. Thus, the observation of iodine cations was also reported more than 80 years ago.\[8\] Meanwhile, several polyiodic cations such as, I$_4$$^-$ and I$_5$$^-$ have been identified, however, again, not as sulfates or polysulfates, but as [AsF$_6$]$^-$ and [SbF$_5$]$^-$ salts, respectively.\[9-13\] The reason for the preference of the anions is, on the one hand that they are weak ligands and on the other, the respective pentafluorides AsF$_5$ and SbF$_5$ are suitable oxidizers which allow oxidation of the heavier chalcogens and halogens in a defined way. Moreover, the pentafluorides can be used in liquid SO$_2$ as a solvent, so that crystallization is facilitated. Another oxidant that has gained popular interest for the preparation of halogen polycations is peroxodisulfuryl difluoride, S$_2$O$_4$F$_2$. After oxidation it provides the [SO$_4$F]$^-$ anion for the stabilization of the generated polycations.\[14\] In the case of iodine the reaction leads to the (I$_4$)$_2$$^2+$ cation, however, some disproportionation to (I$_3$)$_2$$^2+$ and I(SO$_4$F)$_2$ is observed.\[15\] Furthermore, there was clear evidence that the I$_4$$^-$ cation undergoes dimerization under formation of a (I$_4$)$_2$$^2+$ cation at lower temperature in this medium.\[16\] Optically, the dimerization is accompanied by a colour change of the solution from dark blue to red upon cooling. The (I$_4$)$_2$$^2+$ cation has later been isolated in the solid state, namely in form of the salts (I$_4$)[AsF$_6$)$_2$, (I$_4$)[SbF$_6$)$_2$, and (I$_4$)[SbF$_5$][SbF$_6$]$_2$.\[17\] It is noteworthy to mention, that this cation also has not been stabilized in an oxoanionic environment. In general, it can be summarized that the preparation and stabilization of chalcogen and halogen polycations is best achieved using a combination of a sufficiently powerful oxidizer and a weak coordinating anion of a superacid. In retrospect, the initial observations that resulted from experiments in H$_2$SO$_4$ and H$_2$SO$_4$/SO$_3$ mixtures (oleum) are consistent, as these media meet the requirements perfectly. Furthermore, the combination of iodine and oleum (65 % SO$_3$) has been investigated as a potential medium.\[18,19\] The formation of a blue solution has been reported but crystallization attempts were strongly hampered by the viscosity of the solution.

In course of our investigations of the H$_2$SO$_4$/SO$_3$ system we reported several important findings. Firstly, the extraordinary oxidation power could be shown by the oxidation of platinum metals leading to several unique compounds.\[20-23\] Secondly, we could demonstrate that the assumed infinite chain growth of polysulfuric acids H$_n$S$_n$O$_{3n-1}$, that is with increasing SO$_3$ content, is doubtful.\[31,32\] Nevertheless, the assumption that the Bronsted acidity of H$_2$SO$_4$/SO$_3$ mixtures increases with increasing SO$_3$ content seems reasonable.\[10\] Thus, we once more investigated the oxidation of iodine in this medium. In particular, we choose the reaction of I$_3$ with SO$_3$ as the initial experiment. It is known that liquid γ-SO$_3$ (mp. 16.8°C) is extremely prone to polymerization in the presence of water, even at very low H$_2$O concentrations (> 10$^{-4}$ mol %).\[24\] Therefore, the reaction should

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provide a long chained polysulfate anion, as SO₃ is most certainly a sufficiently strong oxidizer.

**Results and Discussion**

The reaction was carried out in such a fashion that elemental iodine and SO₃ were heated to 80 °C in a sealed glass ampoule. Upon cooling to room temperature, a homogeneous turquoise coloured solution was obtained (see Figure S1 in the Supporting Information). After storing the ampoule in a refrigerator at a temperature below 7 °C, a large number of red crystals grew. The agglomerates of these crystals were located next to needle shaped asbestos type SO₃ crystals, which grew simultaneously. Prosaically speaking, the red crystals appear as blossoms at the branches formed by asbestos type SO₃ crystals (Figure 1). The red crystals disappeared if the ampoule is warmed above 7 °C. Moreover, they decompose rapidly after opening of the ampoule, even at low temperature. Thus, the manipulation of the single crystals and their preparation for the measurement has to occur rapidly and under keeping under constant cooling (see Supporting Information). Finally, a single crystal could be measured at the P24 beamline at the DESY facility in Hamburg. The structure analysis revealed that \( (\text{I}_4[\text{S}_6\text{O}_{19}])^{2-} \) has formed.

Neither reducing the amount of SO₃ nor increasing the temperature of the afore introduced mixture led to another product. Thus, with the aim of studying the cation with another anion that still features a polysulfate fragment of the chain, we managed to stabilize the \( (\text{I}_4)^{2+} \) cation with a borosulfate anion, namely the bis-disulfatoborate anion \( [\text{B}((\text{S}_6\text{O}_{19}))_{2}]^{5-} \). As for \( (\text{I}_4[\text{S}_6\text{O}_{19}])^{2-} \), elemental iodine was the starting material and was heated with a mixture of \( \text{SO}_3 \), \( \text{B}_2\text{O}_3 \) and the pyridine-SO₃ complex \( (\text{py-SO}_3) \) to 120 °C. \( (\text{I}_4)[\text{B}((\text{S}_6\text{O}_{19}))_{2}] \) does not crystallize directly from the homogeneous turquoise coloured solution while cooling. However, several days later after having inverted the ampoule, red crystals could be observed on the inner wall (Figure 2).

It is the first time that the \( (\text{I}_4)^{2+} \) cation could be isolated in form of oxaoanionic salts. Interestingly, the stability of the structure results not only from the monovalent anion \( [\text{B}((\text{S}_6\text{O}_{19}))_{2}]^{5-} \) but also thanks to a divalent hexasulfate anion. To date, especially weakly coordinating monovalent anions of superacids were considered as ideal candidates. However, our findings corroborate the above-mentioned theoretical predications, that the acidity of polysulfuric acids \( \text{H}_{2n+1}\text{S}_n\text{O}_{2n+1} \) increases with growing chain lengths. In fact, \( (\text{I}_4)[\text{S}_6\text{O}_{19}] \) is only the third example of a compound containing the hexasulfate anion. The first observations were reported recently with the examples of \( \text{Rb}_2[\text{S}_6\text{O}_{19}] \) and \( (\text{NH}_4)_2[\text{S}_6\text{O}_{19}] \) by ourselves.\(^{[32]}\) Not unknown but also rare is the \( [\text{B}((\text{S}_6\text{O}_{19}))_{2}]^{5-} \) anion. Examples are known, for example, for the alkaline metals and silver in \( \text{M}([\text{B}((\text{S}_6\text{O}_{19}))_{2}]^{-}) \) \( (\text{M} = \text{Li}, \text{Na}, \text{K}, \text{Ag}) \),\(^{[35,36,37]}\) and even more recently we reported on two highly unusual heteropoly Au–Cl cations stabilized by \( [\text{B}((\text{S}_6\text{O}_{19}))_{2}]^{5-} \) anions.\(^{[38]}\)

As described for the fluorides \( (\text{I}_4)[\text{AsF}_{6}]^2- \), \( (\text{I}_4)[\text{SbF}_{6}]^2- \), and \( (\text{I}_4)[\text{SbF}_{6}]^2- \), respectively, the \( (\text{I}_4)^{2+} \) cation in both compounds has a rectangular shape with two short and two longer bonds (Figure 3). In \( (\text{I}_4)[\text{S}_6\text{O}_{19}] \) the short bonds show a value of 260.8(3) pm while the longer edge of the rectangle is 322.1(1) pm long. The values are identical for two of the four bonds because the \( (\text{I}_4)^{2+} \) cation is situated on the 4d site of space group \( \text{C}2/c \), that is, it exhibits inversion symmetry. The respective bond lengths for the three fluorides are smaller for the short edge of the rectangle (\( \alpha = 258 \) pm) and larger for the longer edge (\( \alpha = 327 \) pm). In \( (\text{I}_4)[\text{B}((\text{S}_6\text{O}_{19}))_{2}] \) two opposite bonds are identical and reveal values of 259.2(2) pm. The other two I–I distances are 308.7(2) pm and 339.8(3) pm, meaning they are not identical as in \( (\text{I}_4)[\text{S}_6\text{O}_{19}] \). This finding can be explained by the coordination sphere around the \( (\text{I}_4)^{2+} \) cations, which is very different in the two compounds. Similar to the reported fluorides, there is one shorter contact for each of the iodine atoms to the surrounding anions in \( (\text{I}_4)[\text{S}_6\text{O}_{19}] \). In this case the respective distances I–O are found at 277.8(4) pm and 276.3(4) pm and all others are significantly larger than 300 pm. The situation is different for \( (\text{I}_4)[\text{B}((\text{S}_6\text{O}_{19}))_{2}] \). While the shortest I–O distance for two of the four iodine atoms stems from a \( \mu_2^{-} \)coordinating oxygen atom (I–O: 276.6(3) and 279.6(3) pm), the other two result from a \( \mu_4^{-} \)coordinating oxygen atom (I–O: 294.4(3), 297.3(3) pm). The two
iodine atoms that are coordinated by the μ₂-oxygen atom have two and three additional I–O connections with only little more than 300 pm distance, respectively.

In principle, the formation and the structure of the $\text{(I}_4^+\text{)}^2+$ cation is theoretically well understood: In a first step the oxidation of the I$_2$ molecule leads to shortening of the I–I bond under formation of an $\text{(I}_2^2$+ cation. The latter shows an unpaired electron in a π* orbital and pairing of two of these radicals leads to the $\text{(I}_4^+\text{)}^2+$ ion (see also Supporting material).\(^{[19-41]}\)

$I_4[\text{S}_6\text{O}_{19}]$ is the third example for the existence of the hexasulfate anion. In previous reports we have shown that the stability of polysulfate anions $[\text{S}_n\text{O}_{2n+1}]^{2-}$ against loss of SO$_3$ decreases with growing numbers of $n$. This decreasing stability is strongly correlated with the increasing distance of the terminal S–O bond within the backbone of the anion. The longer the polysulfate chain grows the more the anion can be seen as a Lewis acid/base adduct of the respective smaller cation. The situation for the hexasulfate ion in $\text{(I}_4^+\text{)}[\text{S}_6\text{O}_{19}]$ is significantly different. It is still observed that the terminal sulfur oxygen bond (S3–O231) is the longest within the anion, however, with a value of 177.6(4) pm this bond is dramatically shorter than that found for the rubidium and ammonium compounds, respectively (Figure 4). The difference of more than 40 pm is remarkable and the observed distance is also significantly shorter than the value predicted by theory ($195.3$ pm).\(^{[32,42]}\) Even if the data known so far is scarce, there seems to be clear evidence that the nature of the respective counter cation has a great influence on the bond lengths observed in the hexasulfate anion. The cation in $\text{(I}_4[\text{S}_6\text{O}_{19}]$ is highly unusual compared to the spherical ions used so far. The $\text{(I}_4^+\text{)}^2+$ cation is connected by only two quite short bonds involving the atoms O32 and O33 of the terminal SO$_3$ groups of the $[\text{S}_6\text{O}_{19}]^{2-}$ ion (Figure 4 and 5). We assume that these moieties donate electron density to the π*-orbital of the respective $\text{(I}_4^+\text{)}$ units, causing the significantly different I–I distances of the $\text{(I}_4^+\text{)}^2+$ cation. These strong interactions are also reflected by the bond lengths S3–O32 and S3–O33 (144.6(4) pm and 144.1(4) pm) which are significantly longer than those observed in the hexasulfate anions known so far (~140 pm).

In contrast to $\text{(I}_4[\text{S}_6\text{O}_{19}]$, which exhibits a comparatively symmetrical surrounding of the $\text{(I}_4^+\text{)}^2+$ cation, the coordination sphere of the $\text{(I}_4^+\text{)}^2+$ cation in $\text{(I}_4[\text{B(SO}_3])_2$ is highly unsym-

| Bond | Distance (pm) |
|------|--------------|
| I1–I2 | 259.2(7) |
| I1–I4 | 339.8(3) |
| I1–I3 | 308.7(2) |
| I1–I3 | 259.2(2) |
| I1–O12 | 279.6(3) |
| I1–O12 | 305.0(5) |
| I1–O43 | 363.7(3) |
| I1–O53 | 368.8(3) |
| I1–O61 | 366.9(3) |
| I1–O63 | 345.9(4) |
| I1–O67 | 328.9(3) |
| I1–O73 | 341.5(3) |
| I2–O21 | 364.0(3) |
| I2–O22 | 348.7(4) |
| I2–O22 | 344.7(4) |
| I2–O23 | 354.4(3) |
| I2–O24 | 326.2(4) |
| I2–O52 | 321.9(3) |
| I2–O53 | 350.2(3) |
| I2–O53 | 362.8(4) |
| I2–O72 | 297.3(3) |
| I2–O72 | 277.8(4) |
| I2–O73 | 349.7(3) |
| I2–O73 | 364.0(4) |
| I3–O23 | 368.2(3) |
| I3–O33 | 305.7(3) |
| I3–O33 | 355.5(3) |
| I3–O33 | 345.4(3) |
| I3–O33 | 294.4(3) |
| I3–O33 | 303.7(3) |
| I3–O34 | 349.9(3) |
| I3–O22 | 276.6(3) |
| I4–O22 | 352.3(3) |
| I4–O22 | 358.8(3) |
| I4–O34 | 370.1(3) |
| I4–O62 | 312.5(3) |

Table 1. Selected bond length for $\text{(I}_4[\text{B(SO}_3])_2$ and $\text{(I}_4[\text{S}_6\text{O}_{19}]$.
The different borosulfate anions seem to dictate the structure of the cation. Thus, the longest I–I distance (339.8(3) pm), which is significantly longer than the longest I–I distance in (I$_4$)[B(S$_6$O$_5$)$_2$]$_2$ (322.1(1) pm), is coordinated only by one chelating borosulfate anion with short I–O distances of 259.2(2) and 259.2(7) pm. As for (I$_4$)[S$_6$O$_5$]$_2$ further I–O contacts values > 300 pm can be found. Thus, this finding resembles the situation seen for the hexasulfate anions, in particular exhibiting only chelating μ$_2$-oxygen coordination. However, the opposite side of the (I$_4$)$_2$$^{+}$ cation in (I$_4$)[B(S$_6$O$_5$)$_2$]$_2$ exhibits μ$_2$-oxygen coordination and multiple further contacts to terminal oxygen atoms of borosulfate anions with distances close to 300 pm. Thus, the I–I distance reveals a value of only 308.7(2) pm. The S–O bond lengths of the borosulfate anions are similar to findings for species with spherical monoatomic cations. Thus, the coordination of the borosulfate anion to the (I$_4$)$_2$$^{+}$ cation causes no change in the Lewis acidity of the anion, which is in contrast to the findings for the hexasulfate, where the coordination has a strong influence on the structure of the anion. However, the more compact structure of the borosulfate favours the higher coordination number of the (I$_4$)$_2$$^{+}$ cation and thus influences the bonding situation of the cation significantly.

Conclusions

The successful preparation and characterization of (I$_4$)[S$_6$O$_5$]$_2$ and (I$_4$)[B(S$_6$O$_5$)$_2$]$_2$ provides both, first examples of the (I$_4$)$_2$$^{+}$ cation in an oxoanionic environment and new versions of the rarely seen anions [B(S$_6$O$_5$)$_2$]$^{2-}$ and [S$_6$O$_5$]$_2^{2-}$, displaying remarkable differences with respect to the hexasulfates known so far. Besides these fundamental findings, the investigations illustrate...
once more the enormous potential of \( \text{SO}_3 \) for the preparation of unusual species. In particular, it seems that further scrutiny of non-metallic element reactions in \( \text{SO}_3 \) is worthy of consideration.

### Experimental Section

#### Synthesis

\( \text{SO}_3 \) was obtained in a specially designed apparatus for the generation, distillation and the subsequent transfer into glass ampoules under nitrogen gas. For this purpose, fuming sulfuric acid (5 mL, 65% \( \text{SO}_3 \) used as received, Merck, Darmstadt, Germany) was slowly added via a dropping funnel into a 500 mL flask with \( \text{P}_2\text{O}_5 \) (250 g, > 97%, Merck, Darmstadt). At the same time, the flask was heated at 130 °C and the generated \( \text{SO}_3 \) distilled into a connected burette body (scaling 0.01 ml ± 0.01). A connected glass ampoule \((1 = 200 \text{ mm}, \varnothing = 20 \text{ mm}, \text{ thickness of the tube wall} = 2 \text{ mm})\) containing the solid starting materials was then filled with the suitable amount of \( \text{SO}_3 \). In the case of \((\text{I})[\text{S(O)}_2\text{O}_3]_2\) powder (253 mg, Riedel-de Haën, Seelze) was used and the ampoule then filled with 1.00 mL (24.5 mmol) of freshly distilled \( \text{SO}_3 \). In the case of \((\text{I})[\text{B(S)}_2\text{O}_4])_2\), the pyridine-\( \text{SO}_3 \) complex (284 mg, Sigma Aldrich, Hamburg) was added to \( \text{I}_2 \) powder (150 mg, Riedel-de Haën, Seelze) and \( \text{B}_2\text{O}_3 \) powder (105 mg, Sigma Aldrich, Hamburg). The powders were not ground together in a mortar. The ampoule was then filled with 0.53 mL (14.0 mmol) of freshly distilled \( \text{SO}_3 \). The ampoules were torch sealed and heated in a tube furnace. For the synthesis of \((\text{I})[\text{S(O)}_2\text{O}_3]_2\) a maximum temperature of 80 °C is required, for \((\text{I})[\text{B(S)}_2\text{O}_4])_2\) the reaction temperature is 120 °C. The respective temperatures were maintained for 48 h and the reaction mixtures were then allowed to cool to room temperature at rate of 1/3 °C \(-1\) for \((\text{I})[\text{S(O)}_2\text{O}_3]_2\) and 1 °C \(-1\) for \((\text{I})[\text{B(S)}_2\text{O}_4])_2\), respectively. A homogeneous deep turquoise colored solution was obtained in both cases, from which red crystals crystallized in the refrigerator at 4 °C next to asbestos-like \( \text{SO}_3 \) needles in the case of \((\text{I})[\text{S(O)}_2\text{O}_3]_2\) and after turning the ampoule upside down after several days for \((\text{I})[\text{B(S)}_2\text{O}_4])_2\), besides a colourless by-product. Above 7 °C, the red hexasulfate decomposes within a few minutes and can be re-crystallized again within several days in the refrigerator. The borosulfate is stable at ambient temperature in the closed ampoule. Opening of the ampoule leads to rapid decomposition of the obtained phases. Obviously, the surrounding \( \text{SO}_3 \) atmosphere seems to contribute essentially to the stability of the obtained phases.

**Caution!** \( \text{SO}_3 \) is a strong oxidizer, which needs careful handling. During the reaction and even after cooling down to room temperature the glass tubes may be pressurized. The tubes should be cooled with liquid nitrogen before opening.

#### Structure determination

\((\text{I})[\text{S(O)}_2\text{O}_3]_2\): Crystal structure determination was performed on the P24.1 beamline of the PETRA III facility at German Electron Synchrotron (DESY) Hamburg (Germany). The reaction ampoule containing the red crystalline phase of \((\text{I})[\text{S(O)}_2\text{O}_3]_2\) was stored in the refrigerator at 4 °C until shortly before the measurement. After opening the ampoule, single crystals were transferred to an inert oil with pre-cooled preparation equipment. By continuous cooling in a cold nitrogen stream the individual crystals were separated with the help of a light microscope polarizing filter. Despite the elaborate cooling efforts, the crystals showed spontaneous decomposition on contact with the micromount.

However, due to the surface tension of the inert oil used, a single crystal could be held in the centre of a larger micromount loop without directly touching its edges. Under constant cooling in liquid nitrogen, the attached single crystal was transferred to the cold nitrogen gas stream (100.0(2) K) of the single-crystal diffractometer (Huber 4-circle Kappa, P24 Beamline, Petra III) and intensity data was collected. Later attempts to measure the crystal at higher temperatures led to the loss of the single crystal above 250 K. This observation may be explained by either the temperature or the intensity of the synchrotron radiation used. We favour the first explanation, since all other crystals were likewise decomposed after a few minutes to form a red solution despite cooling under inert oil. We also assume that the obtained crystalline phase of \((\text{I})[\text{S(O)}_2\text{O}_3]_2\) is in fact stabilized within the \( \text{SO}_3 \) atmosphere of the closed ampoule at temperatures < 7 °C. The collected intensity data were reduced, and a cell refinement was carried out.\(^{[45]}\) Intrinsic phasing (SHELXT) gave a successful structure solution.\(^{[46]}\) Finally, anisotropic displacement parameters were introduced, and a multi-scan absorption correction was applied to the reflection data.\(^{[41]}\)

\((\text{I})[\text{B(S)}_2\text{O}_4])_2\): Under a polarization microscope, a suitable crystal was prepared, mounted onto a micromount (MicroMounts™, MSITEgen LLC, New York, USA) and immediately placed into a stream of cold \( \text{N}_2 \) (100(2) K) inside the diffractometer (Bruker Photon III, Bruker, Karlsruhe, Germany). After unit cell determination, the reflection intensities were collected. The collected intensity data were reduced, and a cell refinement was carried out.\(^{[47]}\) The structure solution was successful using intrinsic phasing (SHELXTL).\(^{[41]}\) Finally, anisotropic displacement parameters were introduced, and a multi-scan absorption correction was applied to the reflection data.\(^{[41]}\)

Atomic positions and further details of the crystal structures are given in the Supporting Information. Deposition Numbers \( \text{1955943} \) (for \((\text{I})[\text{S(O)}_2\text{O}_3]_2\)) and \( \text{2149733} \) (for \((\text{I})[\text{B(S)}_2\text{O}_4])_2\)) contain(s) 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.

#### Data Availability Statement

The data that support the findings of this study are available in the supplementary material of this article.
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[1] M. H. Klaproth, Phil. Mag. 1798, 1, 78.
[2] G. Magnus, Ann. Phys. 1827, 10, 491.
[3] D. J. Prince, J. D. Corbett, B. Garbisch, Z. Anorg. Chem. 1970, 9, 2731.
[4] R. K. Mullin, D. J. Price, J. D. Corbett, Z. Anorg. Chem. 1971, 10, 1749.
[5] R. J. Gillespie, J. Passmore, Adv. Inorg. Chem. Radiochem. 1975, 17, 49.
[6] R. J. Gillespie, J. Chem. Soc. 1979, 8, 315.
[7] I. D. Brown, D. B. Crump, R. J. Gillespie, Inorg. Chem. 1971, 10, 2319.
[8] I. Masson, C. Argument, J. Chem. Soc. 1938, 1702.
[9] C. G. Davies, J. R. Gillespie, P. R. Irel, J. M. Sowa, Can. J. Chem. 1974, 52 (11), 2048.
[10] J. Passmore, P. Taylor, T. Whidden, P. S. White, Can. J. Chem. 1979, 57, 968.
[11] J. Passmore, G. Sutherl, P. S. White, Inorg. Chem. 1981, 20(7), 2169.
[12] A. Apblett, F. Grein, J. P. Johnson, J. Passmore, P. S. White, Inorg. Chem. 1986, 25, 422.
[13] J. D. Corbett, Prog. Inorg. Chem. 1976, 21, 129.
[14] C. Chung, G. H. Cady, Inorg. Chem. 1972, 11, 2628.
[15] R. J. Gillespie, K. C. Malhotra, Inorg. Chem. 1969, 8, 1751.
[16] R. J. Gillespie, J. B. Milne, M. J. Morton, Inorg. Chem. 1968, 7, 2221.
[17] R. Faggiani, R. J. Gillespie, R. Kapoor, C. J. L. Lock, J. E. Vekris, Inorg. Chem. 1988, 27, 4350.
[18] W. Gattardi, Mth. Chemie 1975, 106, 1203.
[19] A. Ball, K. C. Malhotra, J. Inorg. Nucl. Chem. 1976, 38, 411.
[20] J. Bruns, T. Klüner, M. S. Wickleder, Angew. Chem. Int. Ed. 2013, 52, 2590; Angew. Chem. 2013, 125, 2650.
[21] J. Bruns, M. Hänsch, M. S. Wickleder, Inorg. Chem. 2015, 54, 5681.
[22] J. Bruns, T. Klüner, M. S. Wickleder, Chem. Asian J. 2014, 9, 1594.
[23] J. Bruns, O. Niehaus, R. Pöttgen, M. S. Wickleder, Chem. Eur. J. 2014, 20, 811.
[24] S. Schwarzer, A. Betke, C. Logemann, M. S. Wickleder, Eur. J. Inorg. Chem. 2017, 3, 752.
[25] M. Pley, M. S. Wickleder, Eur. J. Inorg. Allg. Chem. 2005, 630, 529.
[26] M. Pley, M. S. Wickleder, Angew. Chem. 2004, 116, 4262; Angew. Chem. Int. Ed. 2004, 43, 4168.
[27] A. Arndt, M. S. Wickleder, Z. Anorg. Allg. Chem. 2006, 632, 2104.
[28] M. Pley, M. S. Wickleder, Z. Anorg. Allg. Chem. 2004, 630, 1753.
[29] J. Bruns, D. van Gerven, T. Klüner, M. S. Wickleder, Angew. Chem. 2016, 128, 8253; Angew. Chem. Int. Ed. 2016, 128, 8121.
[30] J. Bruns, M. Eul, R. Pöttgen, M. S. Wickleder, Angew. Chem. Int. Ed. 2012, 51, 2204.
[31] L. V. Schindler, M. Struckmann, A. Becker, M. S. Wickleder, Eur. J. Inorg. Chem. 2017, 5, 958.
[32] L. V. Schindler, A. Becker, M. Wieckhusen, T. Klüner, M. S. Wickleder, Angew. Chem. 2016, 128, 16399; Angew. Chem. Int. Ed. 2016, 55, 16165.
[33] I. A. Koppel, P. Burk, I. Koppel, I. Leito, T. Sonoda, M. Mishima, J. Am. Chem. Soc. 2000, 122, 5114.
[34] R. Grau, W. A. Roth, Z. Anorg. Allg. Chem. 1930, 188, 173.
[35] M. Daub, K. Kasmierzak, P. Gross, H. A. Höppe, H. Hillebrecht, Inorg. Chem. 2013, 52, 6011.
[36] M. Daub, K. Kasmierzak, H. A. Höppe, H. Hillebrecht, Chem. Eur. J. 2013, 19, 16954.
[37] P. Netzsch, H. A. Höppe, Eur. J. Inorg. Chem. 2021, 11, 1065.
[38] S. Sutorius, D. van Gerven, S. Olthof, B. Rasche, J. Bruns, Chem. Eur. J. 2022, 28, e202200004.
[39] J. Passmore, G. Sutherland, T. Whidden, P. S. White, J. Chem. Soc. Chem. Commun. 1982, 289.
[40] M. P. Murchie, J. P. Johnson, J. Passmore, G. W. Sutherland, M. Tajik, T. K. Whidden, P. S. White, F. Grein, Inorg. Chem. 1992, 31, 273.
[41] S. Brownridge, I. Crossig, J. Passmore, H. D. B. Jenkins, H. K. Roobottom, Coord. Chem. Rev. 2000, 197, 397.
[42] C. Logemann, T. Klüner, M. S. Wickleder, Angew. Chem. Int. Ed. 2012, 51, 4997.
[43] Bruker (2018), Apex3. Bruker AXS Inc., Madison, Wisconsin, USA.
[44] G. Sheldrick, Acta Crystallogr. Sect. A 2015, 71, 3.

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