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# $S_6N_2O_{15}$ – a nitrogen-poor sulfur nitride-oxide, *and* the anhydride of nitrido-*tris*-sulfuric acid

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Abstract: The reaction of hexachlorophosphazene,  $P_3N_3Cl_6$ , with SO<sub>3</sub> leads to the new sulfur nitride-oxide  $S_6N_2O_{15}$ . The compound displays an extraordinarily low nitrogen content and exhibits a bicyclic cage structure according to the formulation  $N\{S(O)_2O(O)_2S\}_3N$ , with both nitrogen atoms in trigonal planar coordination of sulfur atoms. Interestingly, the new nitride-oxide can be also seen as the anhydride of nitrido-*tris*-sulfuric acid,  $N(SO_3H)_3$ .

Sulfur trioxide, SO<sub>3</sub>, is an interesting reagent in chemical reactions. On one hand, it is a very strong oxidizer and on the other hand, it can act as a typical Lewis base. We have used the oxidation strength of SO<sub>3</sub>, especially under harsh conditions, for the oxidation of noble metals and noble metal compounds, respectively. The formation of two modifications of  $Pd(S_2O_7)$  by the reaction of elemental palladium with  $SO_3$  is a nice example of these efforts.<sup>[1,2]</sup> On the other hand, SO<sub>3</sub> is a strong Lewis acid and forms readily adducts with several Lewis bases. Well known examples are the complexes with dioxane and pyridine (py).<sup>[3,4]</sup> The latter, SO<sub>3</sub>·py, is even a commercial product that serves as a save SO<sub>3</sub> source for many reactions. In fact, Lewis acid/base adducts with N-donor molecules and SO<sub>3</sub> have been studied quite extensively starting already in the 1950s.<sup>[5]</sup> Actually, very spectacular compounds have been prepared at that time, for example the adducts  $S_4N_4 \cdot xSO_3$  (x = 1-4), for which  $S_4N_4 \cdot SO_3$  has been structurally characterized later.<sup>[6]</sup> Another potential base that has been considered for SO<sub>3</sub> interaction was hexachlorophosphazene, P<sub>3</sub>N<sub>3</sub>Cl<sub>6</sub>.<sup>[5]</sup> It has formally three available nitrogen atoms bearing free electron pairs. Thus, the composition P<sub>3</sub>N<sub>3</sub>Cl<sub>6</sub>·3SO<sub>3</sub> of the reported complex it is very reasonable, even if structural information is still lacking. We came across that compound for two reasons: On one hand, we are interested in Lewis acid/base complexes of SO<sub>3</sub> since we discovered that the rarely known polysulfates  $[S_nO_{3n+1}]^{2-}$  have to be described as adducts according to  $[S_nO_{3n+1}]^{2-}$  SO<sub>3</sub>, at least for larger numbers of *n*.<sup>[7,8]</sup> In the hexasulfate Rb<sub>2</sub>[S<sub>6</sub>O<sub>19</sub>] (*n* = 6), the distance of the sulfur atom of SO<sub>3</sub> to the next oxygen atom is already as long as 231 pm.<sup>[8]</sup> For a detailed investigation of bond lengths within Lewis acid/base complexes structure elucidations of complexes with different bases are desirable. On the other hand, we have recently started a research project aiming at a detailed understanding of nitrogen-based derivatives of sulfuric acid. These are for example the slightly acidic sulfimide,

 $SO_2(NH_2)_2$ ,<sup>[9]</sup> and its cyclic condensation products  $S_3O_6(NH)_3$  and  $S_4O_8(NH)_4$ ,<sup>[10,11]</sup> for which a limited number of salts are known.<sup>[12-14]</sup> However, the more prominent of these derivatives are amidosulfuric acid, imido-*bis*-sulfuric acid, and nitrido-*tris*-sulfuric acid (Fig. 1). Even if all of these acids are textbook examples, our knowledge is still quite limited. Only amidosulfuric acid, in its zwitterionic ground state a Lewis acid/base complex of SO<sub>3</sub> and NH<sub>3</sub>, and amidosulfates have been frequently reported.<sup>[15]</sup> For all of the other anions depicted in figure 1 a very limited number of salts is known.<sup>[16]</sup> Especially for the nitrido-*tris*-sulfuric acid, N(SO<sub>3</sub>H)<sub>3</sub>, which is not known in a pure form, there is just one report of respective salt, namely K<sub>3</sub>[N(SO<sub>3</sub>)<sub>3</sub>]·2H<sub>2</sub>O.<sup>[17]</sup> The acid and their salts are prone to hydrolysis, what is certainly a drawback for synthesis, especially from aqueous solution. In this case, hexachlorophosphazene might be a suitable nitrogen source for the preparation of N-based sulfuric acids under anhydrous conditions.



Figure 1 Nitrogen-based sulfuric acid derivatives

With respect to the two above-mentioned issues, i.e. hexachlorophosphazene as Lewis base and as starting material for the synthesis of N-based sulfuric acid derivates, we have investigated the reaction of SO<sub>3</sub> and P<sub>3</sub>N<sub>3</sub>Cl<sub>6</sub> under various conditions in more detail. According to the findings of Goehring et al., at low temperature ( $\approx 40 \, ^{\circ}$ C) a reaction is observed, however without gaining crystalline material. Only if the temperature is raised up to 80  $^{\circ}$ C a huge number of single crystals growths from excess SO<sub>3</sub> in a short period of time (Fig. 2). Structure elucidation revealed that the anhydride of nitrido-*tris*-sulfuric acid had formed, namely S<sub>6</sub>N<sub>2</sub>O<sub>15</sub>. With respect to the amount of the gained product, the reaction is almost quantitative, so that the reaction could be written as 2 P<sub>3</sub>N<sub>3</sub>Cl<sub>6</sub> + 18 SO<sub>3</sub>  $\rightarrow$  3 S<sub>6</sub>N<sub>2</sub>O<sub>15</sub> + 4 POCl<sub>3</sub> +  $\frac{1}{2}$  P<sub>4</sub>O<sub>10</sub>. We have not identified the by-products unambiguously up to now, however, we do not observe elemental chlorine, which is, according to [5], a reaction product at higher temperature. A very likely product is phosphoryl chloride, POCl<sub>3</sub>. The presence of POCl<sub>3</sub> would also explain that the sulfur trioxide which is used in excess in the reaction stays liquid, even if the ampoules are stored in a refrigerator. In similar reactions, we usually observe the formation of asbestos type sulphur trioxide ( $\alpha$ -SO3) at lower temperature, visible by large needle shaped crystals growing in the ampoule. Compounds like SO<sub>2</sub>Cl<sub>2</sub> or POCl<sub>3</sub> are well-known stabilizers that are used to keep sulphur trioxide liquid below 30 °C by supressing the polymerisation of SO<sub>3</sub> molecules.<sup>[18,19]</sup> Attempts to separate the obtained by-product from SO<sub>3</sub> failed up to now.



Figure 2 Single crystals of S<sub>6</sub>N<sub>2</sub>O<sub>15</sub>

The molecular compound has a unique structure with two three-coordinate nitrogen atoms connected by three  $[S_2O_5]$  groups (Fig. 3), as it would be emphasized by the more descriptive formula N{S(O)<sub>2</sub>O(O)<sub>2</sub>S}<sub>3</sub>N. The distances S-N fall is a narrow range between 170.6 and 171.6 pm, and the surrounding of the nitrogen atoms is almost perfectly planar. Thus, no activity of the lone electron pair is observable, obviously due to significant  $\pi$ -bonding to the sulfur atoms. The observation is in line with the reported findings for the anion  $[N(SO_3)_3]^{3-,[17]}$  The nitrogen atoms are connected by three nearly identical S-O-S bridges, displaying distances 161.8 and 163.8 pm and angles  $\angle$ S-O-S of about 125°. These are the typical values that are, for example, observed for the disulfate ion,  $S_2O_7^{2-}$ . The distances and angles within the  $S_6N_2O_{15}$  molecule are well reflected by quantum mechanical calculations (cf. caption figure 3 and supplement). As expected, the calculations result in C<sub>3h</sub> symmetry for the molecule, while in the solid state (space group C2/c) only C<sub>1</sub> symmetry is found.



**Figure 3** Structure and labelling of the  $S_6N_2O_{15}$  molecule viewed in different directions. The middle picture shows the molecule viewed along an axis through the nitrogen atoms, emphasizing their almost perfect trigonal planar coordination by sulphur atom. At right, the molecule bicyclo[3.3.3]undecane is depicted which represents the  $[S_6O_3N_2]$  cage of  $S_6N_2O_{15}$  (emphasized by black bonds). Selected distances (in pm) compared to theoretical values (in *italics*):  $S(1-6)-O_{terminal}$  (O11, O12; O21, O22; O31, O32; O41, O42; O51, O52; O61, O62)  $\approx$  140.5(2)/141.75, S1-O121 161.8(1)/164.12, S2-O121 163.8(2)/164.2, S3-O341 163.6(2)/164.14, S4-O341 162.3(2)/164.12, S5-O561 163.1(2)/164.13, S6-O561 162.8(2)/164.13, N1-S1 170.6(2)/172.64; N1-S3 171.6(2)/172.67, N1-S5 171.6(2)/172.67, N2-S2 171.5(2)/172.67, N2-S4 171.6(2)/172.69, N2-S6 171.6(2)/172.69.

The core cage of the  $S_6N_2O_{15}$  molecule (emphasized by black bonds in figure 3) has the shape of the bicyclic organic molecule bicyclo[3.3.3]undecane. Such a cage has not been observed before in the chemistry of sulphur nitride-oxides, although a significant number of compounds has been observed in the system S/N/O (Fig. 4).<sup>[20]</sup> With respect to the structural characterizations these compounds show chain structures like  $S_2(NSO)_2^{[21]}$  or  $S_3N_2O_2$ ,<sup>[22,23]</sup> cyclic molecules like  $S_3N_2O_5$ ,<sup>[24,25]</sup>  $S_7N_6O_8$ ,<sup>[26]</sup> and  $S_4N_4O_2$ ,<sup>[27]</sup> as well as ionic species like

 $(NO)_{2}[S_{4}O_{13}]$ .<sup>[28]</sup> The most unique compound among the molecular sulfur nitride-oxides is probably the adduct  $S_{4}N_{4}$ ·SO<sub>3</sub> which has already been mentioned in the introduction.<sup>[6]</sup> Another outstanding molecule is sulfuryl azide,  $SO_{2}(N_{3})_{2}$ ,<sup>[29]</sup> which presents the nitrogen richest molecule in the S/N/O system. The new compound  $S_{6}N_{2}O_{15}$  is up to now the sulfur nitride-oxide with the highest oxygen and the lowest nitrogen content.



Figure 4 Molecular compounds in the system S/N/O according to the atomic ratios

The successful synthesis of  $S_6N_2O_{15}$  by the reaction of  $P_3N_3Cl_6$  and  $SO_3$  leads to several new directions that are worthwhile to be followed. On one hand, the reaction might be also suitable for the preparation of the rarely seen nitrido-*tris*-sulfates, if suitable cations are added to the reaction mixture. On the other hand, variation of the reaction conditions may lead to other species is thinkable, for example the  $[N(SO_3)_2]^{3-}$  ion mentioned in the introduction (cf. Fig. 1). Moreover, even anions with both, *tri*- and *bi*-coordinate nitrogen atoms come into sight, for example the hypothetical anion  $[S_6N_3O_{12}]^{3-}$ . Finally, it is worthwhile to remember that there is no nitridosulfate ion,  $[SN_4]^{6-}$ , known up to now, also not in form of condensed species. This finding for sulfur is in strong contrast to the findings for the neighbor elements silicon and phosphorous. Only in organic derivatives, like the famous  $[S(N^tBu)_4]^{2-}$  ion, a complete nitrogen coordination is possible so far.<sup>[30]</sup>

#### **Experimental Section**

#### Synthesis

SO<sub>3</sub> 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% SO<sub>3</sub>, used as received, Merck, Darmstadt, Germany) was slowly added via a dropping funnel into a 500 mL flask with P<sub>4</sub>O<sub>10</sub> (250 g, > 97%, Merck, Darmstadt). At the same time, the flask was heated at 130 °C and the generated SO<sub>3</sub> distilled into a connected burette body (scaling 0.01 ml  $\pm$  0.01). A connected glass ampoule (*l* = 200 mm,  $\emptyset$  = 20 mm, thickness of the tube wall = 2 mm) containing P<sub>3</sub>N<sub>3</sub>Cl<sub>6</sub> (150 mg, 0.431 mmol) was then filled with 1.92 g (24.0 mmol) of freshly distilled SO<sub>3</sub>. The ampoule was torch sealed and heated to 80 °C in a tube furnace. This temperature was maintained for 48 h and the reaction mixture was then allowed to cool down to room temperature within 100 h. S<sub>6</sub>N<sub>2</sub>O<sub>15</sub> could be obtained in form of transparent highly moisture sensitive crystals, which have to be handled under inert gas conditions.

*Caution!*  $SO_3$  is a strong oxidizer which needs careful handling. During the reaction and even after cooling down to room temperature the glass tube might be under pressure. The tube should be cooled with liquid nitrogen before opened.

#### Structure determination

Crystal structure determination has been performed at the P24.1 beamline of the PETRA III facility at German Electron Synchrotron (DESY) Hamburg (Germany). Therefore, single crystals of  $S_6N_2O_{15}$  were prepared under inert oil and selected with the aid of a polarization filter of a light microscope. Attached to a micromount, one single crystal was positioned in 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. Both the collect intensity data were reduced, and a cell refinement was carried out.<sup>[1]</sup> The structure solution under the assumption of the respective space group was successful by intrinsic phasing (SHELXT).<sup>[2]</sup> Finally, anisotropic displacement parameters were introduced and a multi-scan absorption correction was applied to the reflection data. Atomic positions and further details of the crystal structures can be obtained from the joint CCDC/FIZ Karlsruhe deposition service on quoting the deposition number 1901458.

#### IR analysis

IR spectroscopic data were collected with a Bruker Alpha Platinum spectrometer, using the ATR method (attenuated total reflection) in transmission mode. The spectra were measured in the bulk phase on several collected crystals in the range from 4000 to 570 cm<sup>-1</sup>. The IR spectroscopic data were processed and corrected for atmospheric influence by using the OPUS 7.5 software. Important IR bands in cm<sup>-1</sup> (exptl./calcd.): 1512/1535, 1478/1487, 1232/1253, 889/893, 850/831, 699/708, 531/560, 414/403.

#### Calculation method

A full geometry optimization of the trimer of  $S_6N_2O_{15}$  was performed within density functional theory (DFT) using the PBE0 exchange-correlation functional and a cc-pVTZ basis set. The calculations were also used for assigning the IR frequencies. Throughout the study the Turbomole program package was used.

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