Crystal structure of aminopentamide hydrogen sulfate, \((C_{19}H_{25}N_2O)(HSO_4)\)

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The crystal structure of aminopentamide hydrogen sulfate has been solved and refined using synchrotron X-ray powder diffraction data, and optimized using density functional techniques. Aminopentamide hydrogen sulfate crystallizes in space group \(P2_1/c\) (#14) with \(a = 17.62255(14), b = 6.35534(4), c = 17.82499(10) \text{ Å}, \beta = 96.4005(6)°, V = 1983.906(14) \text{ Å}^3\), and \(Z = 4\). The structure consists of layers parallel to the \(bc\)-plane with hydrogen sulfate anions at the core and aminopentamide cations on the outside. There is a strong charge-assisted \(O49\)\(–H53\)⋯\(O52\) hydrogen bond between the hydrogen sulfate anions. This hydrogen bond links the anions in a chain parallel to the \(b\)-axis. The cation forms a discrete \(N\)\(–H\)⋯\(O\) hydrogen bond to the anion. The amide group also forms two weaker discrete hydrogen bonds to the anion. The three \(N\)\(–H\)⋯\(O\) hydrogen bonds link the cations and anions into columns parallel to the \(b\)-axis. This commercial material from USP contained an unidentified impurity, the powder pattern of which could be indexed on a monoclinic unit cell. The powder pattern has been submitted to ICDD for inclusion in the Powder Diffraction File™ (PDF®).

Key words: aminopentamide, Centrine®, powder diffraction, Rietveld refinement, density functional theory

I. INTRODUCTION

“Aminopentamide sulfate” (sold under the brand name Centrine®) is used to control vomiting, diarrhea, and gastrointestinal (GI) pain or spasms in dogs and cats. The observed anti-spasmodic properties of aminopentamide hydrogen sulfate can help control the discomfort associated with anorectal disease. The systematic name (CAS Registry Number 20701-77-3) is 4-(dimethylammonium)-2,2-diphenylpentanamide hydrogen sulfate. A two-dimensional molecular diagram is shown in Figure 1.

The synthesis of aminopentamide was claimed in US Patent 2,647,926 (Speeter, 1953; Bristol Laboratories). The use of aminopentamide hydrogen sulfate in an anesthetic compound was claimed in US Patent 3,896,221 (Christie and Buckwalter, 1975; Bristol-Myers). The pharmacological activity of Centrine was reported in Hoekstra et al. (1954). We are unaware of any published powder diffraction data on aminopentamide hydrogen sulfate.

This work was carried out as part of a project (Kaduk et al., 2014) to determine the crystal structures of large-volume commercial pharmaceuticals and include high-quality powder diffraction data for them in the Powder Diffraction File (Gates-Rector and Blanton, 2019).

II. EXPERIMENTAL

Aminopentamide hydrogen sulfate was a commercial reagent, purchased from USP (Batch #F1B273), and was used as-received. The white powder was packed into a 1.5 mm diameter Kapton capillary and rotated during the measurement at ∼50 Hz. The powder pattern was measured at 295 K at beamline 11-BM (Antao et al., 2008; Lee et al., 2008; Wang et al., 2008) of the Advanced Photon Source at Argonne National Laboratory using a wavelength of 0.458208(2) Å from 0.5 to 50° \(2\theta\) with a step size of 0.001° and a counting time of 0.1 s per step. The high-resolution powder diffraction data were collected using twelve silicon
crystal analyzers that allow for high angular resolution, high precision, and accurate peak positions. A silicon (NIST SRM 640c) and alumina (SRM 676a) standard (ratio Al₂O₃: Si = 2:1 by weight) was used to calibrate the instrument and refine the monochromatic wavelength used in the experiment.

The pattern was difficult to index, until we allowed the possibility of up to 3 unindexed lines among the 20 peaks \((I_{rel} > 1\%\) using DICVOL14 (Louër and Boulitif, 2014). A primitive monoclinic cell with \(a = 17.7176\), \(b = 6.3740\), \(c = 17.9396\) Å, \(\beta = 96.537^\circ\), \(V = 2012.77\) Å³, and \(Z = 4\) was obtained, with figures of merit \(M(18)\) (de Wolff, 1968) and \(P(18)\) (Smith and Snyder, 1979) = 20.7 and 178.6. A reduced cell search in the Cambridge Structural Database (Groom et al., 2016) yielded five hits, but no structures of aminopentamide derivatives. Neither a name search on “aminopentamide” in the PDF-4 Organics database, nor a traditional search/match on the powder pattern yielded any hits.

An aminopentamide molecule was downloaded as Conformer3D_CID_22565.sdf from PubChem (Kim et al., 2019). The file was converted to a *.mol2 file using Mercury (Macrae et al., 2020) and to a Fenske-Hall Z-matrix using OpenBabel (O’Boyle et al., 2011). The structure was solved using Monte Carlo simulated annealing (parallel tempering) techniques as implemented in FOX (Favre-Nicolin and Černý, 2002), with \((\sin\theta/\lambda)_{\text{max}} = 0.3\) Å⁻¹. The suggested space group was \(P2_1/c\), which was confirmed by successful solution and refinement of the structure. Analysis of potential hydrogen bonding interactions indicated that N2 was protonated (N2⋯O50 = 2.612 Å), so H47 was added to it. The O49⋯O52 distance of 2.876 Å indicated that a proton was between these two atoms, so for the refinement H53 was placed at the midpoint of this interatomic vector. Twenty-two “extra” peaks could be indexed by DICVOL14 on a primitive monoclinic cell with \(a = 17.7067\), \(b = 4.1812\), \(c = 38.2550\) Å, \(\beta = 94.907^\circ\), and \(V = 2821.82\) Å³. This phase was added to the refinement as a second Le Bail phase. This Le Bail phase accounts for the non-overlapped impurity peaks, but has the potential to distort the intensities of the major phase in regions where there is peak overlap. Without a structure model or a pure sample of this phase, its concentration cannot be quantified without the addition of an internal standard.

Rietveld refinement was carried out using GSAS-II (Toby and Von Dreele, 2013). Only the 1.4–25.0° portion of the pattern was included in the refinement (\(d_{\min} = 1.058\) Å). The region 1.60–1.95° 2\(\theta\), which contains scatter from the Kapton capillary, was excluded. All non-H bond distances and angles were subjected to restraints, based on a Mercury/Mogul Geometry Check (Bruno et al., 2004; Sykes et al., 2011). The Mogul average and standard deviation for each quantity were used as the restraint parameters. The restraints contributed 1.8% to the final \(\chi^2\). The hydrogen atoms were included in calculated positions, which were recalculated during the refinement using Materials Studio (Dassault Systèmes, 2021). The \(U_{iso}\) of the heavy atoms were grouped by chemical similarity. The \(U_{iso}\) for the H atoms were fixed at 1.3\(x\) the \(U_{iso}\) of the heavy atoms to which they are attached. The peak profiles were described using the generalized microstrain model. The background was modeled using a 6-term shifted Chebyshev polynomial, and a peak at 6.06° 2\(\theta\) to model the scattering from the Kapton capillary and any amorphous component.

The final refinement of 110 variables using 23 279 observations and 75 restraints yielded the residuals \(R_w = 0.1161\) and \(GOF = 2.02\). The largest peak (at S48) and hole (1.37 Å from H44) in the difference Fourier map were \(0.30(7)\) and \(-0.31(7)\) eÅ⁻³, respectively. The largest errors in the difference plot (Figure 2) are in the shapes and intensities of some of the strong low-angle peaks.

The crystal structure was optimized using VASP (Kresse and Furthmüller, 1996) (fixed experimental unit cell) through the MedeA graphical interface (Materials Design, 2016).
calculation was carried out on 16 2.4 GHz processors (each with 4 GB RAM) of a 64-processor HP Proliant DL580 Generation 7 Linux cluster at North Central College. The calculation used the GGA-PBE functional, a plane wave cutoff energy of 400.0 eV, and a $k$-point spacing of 0.5 Å$^{-1}$ leading to a $1 \times 2 \times 1$ mesh, and took $\sim$42 h. A single-point density functional calculation (fixed experimental cell) and population analysis were carried out using CRYSTAL17 (Dovesi et al., 2018). The basis sets for the H, C, N, and O atoms in the calculation were those of Gatti et al. (1994), and that for S was that of Peintinger et al. (2013). The calculations were run on a 3.5 GHz PC using 8 $k$-points and the B3LYP functional, and took $\sim$3.3 h.

III. RESULTS AND DISCUSSION

The root-mean-square Cartesian displacement between the Rietveld-refined and DFT-optimized structures of the aminopentamide cation is 0.361 Å (Figure 3); the maximum difference is 0.769 Å at the methyl group C16. The agreement is at the upper end of the range of correct structures (van de Streek and Neumann, 2014). The agreement of the absolute positions of the cations and anions in the unit cell is reasonable (Figure 4). This discussion concentrates on the DFT-optimized structure. The asymmetric unit (with atom numbering) is illustrated in Figure 5. The best view of the crystal structure is down the $b$-axis (Figure 6). The structure consists of layers parallel to the $bc$-plane with hydrogen sulfate anions at the core and aminopentamide cations on the outside. The interlayer regions consist of parallel and herringbone stacking of phenyl rings.

All of the bond distances and angles fall within the normal ranges indicated by a Mercury/Mogul Geometry check (Macrae et al., 2020). The torsion angle C9–C4–C5–C6 is flagged as unusual; this lies on the tail of a major gauche population of similar torsion angles. The unusual O1–C9–C4–C7 and C7–C4–C9–N3 torsion angles lie in a broad distribution of a small number of similar torsion angles. The C10–C6–C5–C4 torsion angle lies in a minor gauche population of mainly trans torsion angles. The conformation of the cation is slightly unusual.

Quantum chemical geometry optimization of the aminopentamide cation (DFT/B3LYP/6-31G*/*water) using Spartan ’18 (Wavefunction, Inc., 2020) indicated that the observed conformation is 10.2 kcal mol$^{-1}$ higher in energy than the local minimum (Figure 7). A conformational analysis (MMFF force field) indicates that the minimum-energy conformation is 7.2 kcal mol$^{-1}$ lower in energy; the conformational differences are spread throughout the cation (Figure 8). Intermolecular interactions thus affect the solid-state conformation.

Analysis of the contributions to the total crystal energy of the structure using the Forcite module of Materials Studio (Dassault Systèmes, 2021) suggests that the intramolecular deformation energy is dominated by angle deformation.
terms. The intermolecular energy is dominated by electrostatic attractions, which in this force field analysis also include hydrogen bonds. The hydrogen bonds are better analyzed using the results of the DFT calculation.

Hydrogen bonds are prominent in the structure (Table I). There is a strong charge-assisted O49–H53⋯O52 hydrogen bond between the hydrogen sulfate anions. The energy of this hydrogen bond was calculated using the correlation of Rammohan and Kaduk (2018). This hydrogen bond links the anions parallel to the \( b \)-axis in a chain with a graph set \( C1,1(4) \). There is also a weaker O49–H53⋯S48 interaction. The cation forms a discrete N2–H47⋯O50 hydrogen bond to the anion. The energies of the N–H⋯O hydrogen bonds were calculated using the correlation of Wheatley and Kaduk (2019). The amide group N3/H39/H40 also forms two weaker discrete hydrogen bonds to the anion. The three N–H⋯O hydrogen bonds link the cations and anions into columns parallel to the \( b \)-axis. There are three C–H⋯O hydrogen bonds, both to the anion and to the carbonyl oxygen O1. The

Figure 6. The crystal structure of aminopentamide hydrogen sulfate, viewed down the \( b \)-axis. Image generated using diamond (Crystal Impact, 2022).

Figure 7. Comparison of the VASP-optimized (blue) and local minimum-energy conformations (orange) of the aminopentamide cation.

Figure 8. Comparison of the VASP-optimized (blue) and global minimum-energy conformations (green) of the aminopentamide cation.
The preferred orientation was slight in this rotated capillary. A second-order spherical harmonic model was included in the refinement. The texture index was 1.014(0), indicating that preferred orientation was slight in this rotated capillary.

**TABLE I.** Hydrogen bonds (CRYSTAL17) in aminopentamide hydrogen sulfate.

| H-Bond       | D-H (Å) | H···A (Å) | D···A (Å) | D-H···A (°) | Overlap (e) | E (kcal mol⁻¹) |
|--------------|---------|-----------|-----------|------------|-------------|---------------|
| O49–H53···O52| 1.041   | 1.530     | 2.557     | 167.8      | 0.090       | 16.4          |
| O49–H53···S48| 1.041   | 2.241     | 3.299     | 134.9      | 0.012       |               |
| N2–H47···O50 | 1.067   | 1.600     | 2.662     | 172.5      | 0.085       | 6.7           |
| N3–H39···O49 | 1.029   | 1.935     | 2.964     | 177.8      | 0.040       | 4.6           |
| N3–H40···O50 | 1.014   | 2.450     | 2.844     | 102.3      | 0.009       | 2.2           |
| C5–H23···O52 | 1.102   | 2.726     | 3.770     | 158.0      | 0.013       |               |
| C15–H33···O51| 1.096   | 2.603     | 3.595     | 150.2      | 0.014       |               |
| C16–H37···O1 | 1.096   | 2.647     | 3.689     | 158.6      | 0.012       |               |
| C10–H28···C14| 1.096   | 2.440     | 3.339     | 138.4      | 0.013       |               |
| C21–H45···C18| 1.089   | 2.460     | 3.473     | 151.2      | 0.010       |               |

*aIntramolecular.*

Figure 9. The Hirshfeld surface of aminopentamide hydrogen sulfate. Intermolecular contacts longer than the sums of the van der Waals radii are colored blue, and contacts shorter than the sums of the radii are colored red. Contacts equal to the sums of radii are white.

The Bravais–Friedel–Donnay–Harker (Bravais, 1866; Friedel, 1907; Donnay and Harker, 1937) morphology suggests that we might expect platy morphology for aminopentamide hydrogen sulfate, with {100} as the major faces. A second-order spherical harmonic model was included in the refinement. The texture index was 1.014(0), indicating that preferred orientation was slight in this rotated capillary.

The authors have no conflicts of interest to declare.

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**CONFLICT OF INTEREST**

The authors have no conflicts of interest to declare.
