Original paper

Crystal structure of the uranyl–molybdate mineral calcurmolite Ca\([(\text{UO}_2)_3(\text{MoO}_4)_2(\text{OH})_4](\text{H}_2\text{O})_{-5.0}; insights from a precession electron-diffraction tomography study

Gwladys STECIUK¹*, Radek ŠKODA², Jan ROHLÍČEK¹, Jakub PLÁŠIL¹

¹ Institute of Physics, Academy of Sciences of the Czech Republic v.v.i, Na Slovance 2, 182 21 Prague 8, Czech Republic; steciuk@fzu.cz
² Department of Geological Sciences, Masaryk University, Kotlářská 2, 611 37 Brno, Czech Republic

Calcurmolite is a rare supergene U mineral formed during the alteration–hydration weathering of uraninite and hypogene Mo minerals; its structure has remained unsolved owing to a lack of crystal material suitable for conventional structure analysis. Here, single-crystal precession electron-diffraction tomography shows the calcurmolite (Rabejac, France) structure to be modulated; it is triclinic, crystallizing in the super-space group $P1(a000)$, with $a = 3.938 \text{ Å}$, $b = 11.26 \text{ Å}$, $c = 14.195 \text{ Å}$, $\alpha = 84.4^\circ$, $\beta = 112.5^\circ$, $\gamma = 133.95^\circ$ and has a modulation vector $q = 0.4 a^\ast$. Due to the poor quality of diffraction data, only a kinematical refinement was undertaken, although final results were reasonable: $R\_all = 0.3825/0.3834$ for 3953/17442 observed/all reflections. The structure of calcurmolite is based upon the infinite uranyl–molybdate sheets with baumoite topology ($U : Mo$ ratio = 1.5) and an interlayer of 6-coordinated Ca$^{2+}$ cations with interstitial $\text{H}_2\text{O}$. Adjacent sheets are linked via Ca–O, as well as H-bonds. The structure formula, based on assumed occupancies in the supercell $5a \times b \times c$, is $\text{Ca}(\text{UO}_2)_3(\text{MoO}_4)_2(\text{OH})_4(\text{H}_2\text{O})_{-5.0}$ (for $Z = 4$).

Keywords: calcurmolite, uranyl–molybdate, crystal structure, modulation, baumoite topology, precession electron diffraction

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1. Introduction

Uranyl–molybdates are uncommon alteration products that result from the weathering of uraninite $\text{UO}_2^{2-}$, associated with some hypogene molybdenum-containing minerals, such as molybdénite $\text{MoS}_2$, or its Fe-rich variety, called ‘femolite’ (Mo,Fe)$\text{S}_2$. The latter femolite is less stable under oxidizing conditions (Krivovichev and Plášil 2013). Currently, only nine uranyl–molybdates are known from oxidized parts of uranium deposits worldwide; nevertheless, their crystal chemistry is particularly interesting as it diverges from other known $\text{U}^{6+}$-containing compounds (Burns 2005). The family of synthetic molybdates is considerably broad (e.g. Krivovichev and Burns 2000a, b, 2001a, b, 2002a, b, c, d; Krivovichev et al. 2002); it may be speculated that some of these will be discovered in nature.

Originally, calcurmolite was described by Rudnitskaya (1959) as a new mineral from the Kadjaran ore field, the Kafan region, Armenia, under the name ‘kadjaranite’. Later on, it was found by Fedorov (1963) and Skvortsova et al. (1969) in Kazakhstan at U–Mo deposits, and then reported by Deliens (1992) from Rabejac, France. Sidorenko et al. (2005) presented new chemical data from electron microprobe, infrared (IR) spectroscopy and powder X-ray diffraction for samples from Kazakhstan and Armenia. They demonstrated that calcurmolite forms fine intergrowths with uranophane and uranocircite, which makes its detailed study difficult. On the basis of IR spectra, Sidorenko et al. (2005) concluded that, similar to umohoite, iriginite and mourite, calcurmolite does not contain $\text{(MoO}_4)^{2-}$ tetrahedral oxyanions, but $\text{Mo}^{6+}$ appears in octahedral coordination. This conclusion was confirmed by Frost et al. (2008) by Raman spectroscopy studies of calcurmolite. Calcurmolite minerals of different origins were already reported as poorly crystallized phases by Deliens (1992) and Dal Bo (2018). Up to now, reliable structure data were lacking for this mineral and the chemical formula represented only an uncertain estimate.

Here we report the crystal structure of calcurmolite for the first time based on a precession electron-diffraction tomography study carried out on a calcurmolite sample from the Lodyève Basin in France. This method is a good alternative to single-crystal X-ray diffraction for samples like calcurmolite made of small crystals ($c. 1–5 \text{ µm}$). Moreover, despite the generally poor crystallization ability of the mineral, small parts of crystals giving satisfying diffraction signal were found at the nanoscale allowing a structural determination.
2. Occurrence

The specimen of calcurmolite investigated here originates from the Rabejac deposit in the Lodève Basin, located in the southern part of Massif Central, France. More detailed information about the locality, and its mineralogy and geology, can be found in Dal Bo (2018). Calcurmolite is present as bright yellow nodules resembling mimosa flowers constituted by a radial aggregation of tiny prismatic crystals with a section of 1–5 µm (Fig. 1). In the studied sample, calcurmolite is associated with reddish-brown umohoite and pale yellow uranophane-β. The sample matrix consists of a fine-grained sandstone with the uranyl minerals mentioned above being scattered on the surface.

3. Chemical composition

The chemical composition of calcurmolite was determined using a Cameca SX 100 electron microprobe operated in WDS mode. An operating voltage of 15 kV, beam current of 10 nA and a 10 µm beam diameter and the following X-ray lines and standards were used: \( K_a \) lines: Na (albite), K (orthoclase), Ca (andradite), Al (gahnite), V (vanadinite), Fe (almandine); \( L_a \) lines: Mo (metallic Mo), As (lammerite); \( M_a \) lines: Pb (vanadinite); \( M_b \) lines: U (UO\(_2\)). Other elements were sought but not detected. Peak counting times were 10–20 s and the counting time for the background was 50 % of that of the peak. Matrix effects were accounted for using the \( PAP \) correction routine (Pouchou and Pichoir 1985). Because of the paucity of pure material required for direct \( \text{H}_2\text{O} \) determination, molecular \( \text{H}_2\text{O} \) was calculated based on stoichiometric constrains. Chemical microanalysis revealed the rather homogeneous nature of the specimen. The empirical formula, calculated as the mean of 13 representative points (Tab. 1), is

\[
(K_{0.005}Na_{0.005})\Sigma_{0.01}Ca_{0.16}\left[(UO_2)_{0.6}O_{1.6}Mo_{0.395}(OH)_{0.80}\right](H_2O)_{~1.0}
\]

based on 0.6 \( \text{U} \ apfu \), stoichiometry derived from the modulated structure with \( Z = 4 \). In the \( 5a \times b \times c \) supercell, the composition becomes

\[
(K_{0.025}Na_{0.025})\Sigma_{0.05}Ca_{0.8}[(UO_2)_3O_8Mo_{1.975}(OH)_4](H_2O)_{~5.0}
\]

for \( Z = 4 \). The ideal chemical formula in the supercell is \( \text{Ca}[(UO_2)_3(MoO_4)_2(OH)_4](H_2O)_{~5} \).

4. Structure determination

4.1. Powder X-ray diffraction (PXRD)

For the collection of powder X-ray diffraction data, the sample was ground and placed to the 0.3 mm borosilicate-glass capillary. Powder diffraction data were collected in Debye–Scherrer transmission at

Tab. 1 Chemical composition of calcurmolite from WDS (in wt. % oxides)

|       | Mean (\( n = 13 \)) | Range       | SD   |
|-------|---------------------|-------------|------|
| CaO   | 3.49                | 3.37–3.63   | 0.08 |
| Na\(_2\)O | 0.04              | 0.00–0.12   | 0.05 |
| K\(_2\)O | 0.13               | 0.00–0.26   | 0.09 |
| MoO\(_3\) | 22.17             | 21.10–23.11 | 0.57 |
| UO\(_2\) | 67.18             | 64.48–68.13 | 0.90 |
| H\(_2\)O (calc.) | 9.94*             |             |      |
| Total | 102.50             |             |      |

* calculated from the structure

SD – standard deviation
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150 K on a Empyrean of PANalytical (λ (CuK\text{α}) = 1.54184 Å) powder diffractometer, equipped with the focusing mirror, capillary holder and PIXcel\textsuperscript{3D} detector. The most characteristic calcurmolite peaks are found for interplanar distances of 9.115, 8.03, 7.746, 4.01, and 3.876 Å (Fig. 2). The powder X-ray diffraction pattern shows a lot of similarities with previous study of calcurmolite from Rabejac (Deliens 1992) and from Kazakhstan and Armenia (Sidorenko et al. 2005), but cannot be considered as being identical. The strongest similarities are found with the second sample from Kazakhstan (Sidorenko et al. 2005) where all the characteristic peaks mentioned above are present with comparable relative intensities. This observation is surprising as the composition of calcurmolite from Rabejac in this study is closer to other sample of the same origin (Deliens 1992) than calcurmolite from Kazakhstan (Sidorenko et al. 2005).

The X-ray powder pattern did not provide any further, useful structural information due to the poorly crystalline nature of calcurmolite as well as the presence of significant quantities of uranophane-β. Though, the powder X-ray diffraction data were used to refine the unit cell parameters determined by electron diffraction (see later) and to validate that the crystals measured at the nanoscale were representative of the mineral.

4.2. Precession electron-diffraction tomography (PEDT)

As the low crystallinity and the size of calcurmolite crystals prevented single-crystal X-ray structure studies (all performed attempts failed), structure data were obtained from electron-diffraction and collected with a FEI Tecnai 20 transmission-electron microscope (TEM, acceleration voltage of 200 kV, LaB\text{6}) using a Gatan cryo-transfer holder. For TEM investigations, a small portion of calcurmolite crystals was mildly crushed without solvent and deposited on a Cu-grid with a thin film of holey amorphous carbon. The grid was plunged into liquid nitrogen and then transferred to the TEM. Precession electron-diffraction tomography (PEDT) data sets of non-oriented patterns were collected at 100 K on several crystals (eight data sets) with the precession device Nanomegas Digistar and a side-mounted CCD camera Olympus Veleta with 14-bit dynamic range (Fig. 3a). The precession angle was set to 1° and a tilt step to 1° as well. A condenser aperture of 10 mm and a low illumination setting (spot size 8) were used to reduce the electron dose.

The PEDT data sets were analysed using programs PETS2.0 (Palatinus et al. 2019) and Jana2006 (Petticke et al. 2014). Albeit the crystals were poorly crystallized, at the nanoscale, a few better-crystallized areas of crystals...
were discovered making structural analysis possible by combining the two best data sets (Tab. 2). Only two of the eight measured data sets were used in the structural analysis because of limitations present in PEDT data. First, despite that the ordering along \( a^* \) is well visible on \( h0lm \), as soon as the crystal was rotated during the PEDT experiments, diffuse scattering appeared along the stacking direction \( b^* \) (Figs 3b and 4b) and the resolution was significantly lowered. Moreover, because of the preferred orientation of needle-like crystals on the flat grid (Fig. 3a), the data completeness is limited to 76% for \( \sin \theta/\lambda = 0.7 \ \text{Å}^{-1} \) resolution. Combining more data sets (four were of acceptable quality) did not improve the data completeness and drastically increased the \( R_{int} \) value.

The reciprocal space was first indexed using the superspace formalism in a standard triclinic unit cell: \( a = 3.938 \ \text{Å}, b = 8.15 \ \text{Å}, c = 13.14 \ \text{Å}, \alpha = 104.7^\circ, \beta = 96.4^\circ, \gamma \approx 90^\circ, V = 409.7 \ \text{Å}^3 \) and a modulation vector \( q = 0.4 \ a^* + 0.8 \ b^* + 0.4c^* \) with satellite reflections up to 2\textsuperscript{nd} order (Fig. 4a). In Fig. 4a the reciprocal space is projected into one unit cell to show that, despite the presence of diffuse features and the signal coming from other orientations, most of the reflections are indexed. However, because the completeness is low and the structure needs to be manually interpreted layer by layer, a non-standard choice was preferred with the following parameters: \( a = 3.938 \ \text{Å}, b = 11.26 \ \text{Å}, c = 14.195 \ \text{Å}, \alpha = 84.4^\circ, \beta = 112.5^\circ, \gamma = 133.95^\circ \) and a modulation vector \( q = 0.4 \ a^* \) with satellite reflections up to 2\textsuperscript{nd} order (\( V = 409.72 \ \text{Å}^3 \)) (Fig. 4a). With these settings the layers can be represented in the supercell \( 5a \times b \times c \) in \( (a, c) \) planes, the modulation occurs along \( a^* \) and the layers are stacked following the \( b \)-axis. Based on few peaks from PXRD patterns, Dal Bo (2018), Deliens (1992) and Sidorenko et al. (2005) sug-
gested that calcurmolite could be indexed in a monoclinic or pseudo-orthorhombic unit cell. From PEDT single-crystal data, only triclinic unit cells are found. The intensity integration was performed using the new options available in PETS2.0 using the non-standard settings. The experimental intensity profile (rocking curve) is fitted using several parameters including the rocking curve width (RC width = 0.017 Å⁻¹) and the apparent mosaicity (mosaicity = 0.25°) (for detailed information, see Palatinus et al. 2019). The values of the apparent mosaicity and RC width are very high for calcurmolite data because of the presence of the disorder features along b* and the high mosaicity of the crystals (for well crystallized crystals measured with the same microscope: RC width ≤ 0.003 Å⁻¹ and mosaicity ≤ 0.05°).

The structure was solved using the charge-flipping algorithm (Palatinus and Chapuis 2007) in the superspace group (SSG) P-1(a00)0 and later refined in the SSG P1(a00)0. Because of all the limitations presented previously, the structure solution did not arise directly from the charge-flipping algorithm; the initial result, a 3D electrostatic potential map (e-map) represented as isosurfaces, had to be interpreted manually (Fig. 5). The e-map shows a structure resulting from the alternation of two different layers: the first having strong electrostatic potentials attributed to a uranium–molybdenum layer: the second containing calcium and H₂O that is not resolved enough to be interpreted directly from the e-map (Fig. 5a).

In the U–Mo layer (Fig. 5b), by lowering the isosurfaces level of the potential map to 1σ[ΔV(r)], the visible oxygen positions define two possible coordination polyhedra around the cations: an octahedral coordination and a pentagonal bipyramid coordination. These are characteristic of MoVI(MoV) and U(VI) environments, respectively (Fig. 5b). Following these observations, in the modulated description (SSG P-1(a00)0), Mo and U share the same two cationic sites and their ordering is described using a discontinuous crenel-like modulation function with Δ (U1,2) = 0.6 and Δ (Mo1,2) = 0.4, the width of the crenels (crenel-like; Petříček et al. 2016) associated with one harmonic function. The apical oxygen sites (Oa1, Oa2, Ob1 and Ob2) functions as well as the O3 and O4 atoms located within the U–Mo plane are defined without modulation due to lack of information from the data. The additional oxygen sites O5a, O5b, O5c and O5d of the U–Mo plane are described using crenel functions. The position of the atomic sites described with crenel functions as a function of the α component of the modulation vector shows that the origin of the modulation is explained by the ordering of uranium and molybdenum along a*.

The refinement of the initial model was performed in the final superspace group P1(a00)0 using the kinematical approximation against PEDT data in order to reveal the topology of the Ca–O layer. In the case of calcurmolite, the poor quality of the data as well as the large number of refined parameters prevented the use of the dynamical refinement (Palatinus et al. 2015a, b). Indeed, the presence of disorder reduces the dynamical effects and hence makes refinement using the kinematical approximation relevant. Distance restrictions were applied to maintain the polyhedra of the U–Mo layer. The calcium-based layer was revealed from the difference–Fourier map, with two Ca sites in the 6- and 7-coordination, and four anion sites (occupied by molecular water) (Fig. 7). The occupancies of the Ca sites

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**Tab. 2 Summary of data collection conditions and refinement parameters for calcurmolite**

| Structural formula in the modulated cell | Structural formula in the supercell* |
|-----------------------------------------|--------------------------------------|
| Ca₆[(UO₂)₉O₆Mo₆(OH)₈]((H₂O)₆•H₂O) | Ca₆[(UO₂)₉O₆Mo₆(OH)₈]((H₂O)₆•H₂O) |

| Unit-cell parameters (PEDT) | a = 3.938 Å, b = 11.26 Å, c = 14.195 Å, α = 84.4°, β = 112.5°, γ = 133.95°, q = 0.4 a* |
|-----------------------------|--------------------------------------------------------------------------------------------------|
| At 100 K                    |--------------------------------------------------------------------------------------------------|
| Unit-cell parameters (PXRD) | a = 3.927(2) Å, b = 11.500(4) Å, c = 14.175(2) Å, α = 85.20(2)°, β = 111.70(2)°, γ = 134.05(2)°, q = 0.4 a* |
| At 150 K                    |--------------------------------------------------------------------------------------------------|
| V (100K)                    | 409.7(2) Å³                                                                                     |
| Z                           | 4                                                                                                |
| Density [g.cm⁻³] (from PEDT) | 4.2124                                                                                          |
| Space group                 | P1(a00)0                                                                                        |
| Temperature                 | 100 K                                                                                            |
| Transmission electron microscope | FEI Tecnai 02                                       |
| Radiation (wavelength)      | electrons, (0.0251 Å)                                                                           |
| Resolution                  | 0.1–0.7 Å⁻¹                                                                                     |
| Limiting Miller indices     | –6 < h < 6, –15 < k < 15, –19 < l <19, –2 < m < 2 all : 3953/17442                           |
| No. of independent reflections (obs/all) – kinematic | 1343/3501                                                                                     |
|                            | order 1 : 1736/6993 order 2 : 874/6998                                                          |
| Coverage for sinθ/λ = 0.7 Å⁻¹ | 76.06 %                                                                                         |
| Rw(obs/all) – kinematic     | 0.2374/0.3901                                                                                    |
| Redundancy                  | 1.715                                                                                            |
| R, wR (obs) (kinematical)   | all : 0.3825/0.3834 main : 0.3313/0.3445 order 1 : 0.3887/0.4136 order 2 : 0.5441/0.6170       |

N refined parameters = 100

*5a×b×c supercell
Fig. 4a – Projection of the reciprocal space into one modulated unit cell and two orders of satellite reflections for the standard settings and the settings used in this work. b – Two reciprocal space sections showing the ordering along the $a$ axis on $h0lm$ and the low resolution together with the disorder along the stacking direction $b^*$ visible on $hk0m$.

Fig. 5a – Isosurfaces of the 3D electrostatic potential map (e-map) extended along the stacking direction. Isosurface levels are $2\sigma[\Delta V(r)]$. b – [010]* e-map projection of the layer (1σ[ΔV(r)]) containing uranium, molybdenum and oxygen with its interpretation. The layer model is represented in the supercell $5a \times c$. c – Coordination polyhedra of U$^{6+}$ and Mo$^{6+}$. 

| a) Standard settings | b) Projection of the reciprocal space into one modulated unit cell and two orders of satellite reflections for the standard settings and the settings used in this work. | b) Two reciprocal space sections showing the ordering along the $a$ axis on $h0lm$ and the low resolution together with the disorder along the stacking direction $b^*$ visible on $hk0m$. |
|----------------------|---------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------|
| $a = 3.938 \ \text{Å}, b = 8.15 \ \text{Å}, c = 13.14 \ \text{Å}$, $\alpha = 104.7^\circ$, $\beta = 96.4^\circ$, $\gamma = 90^\circ$, $q = 0.4a^* + 0.8b^* + 0.4c^*$, $V = 409.7 \ \text{Å}$. | $a = 3.938 \ \text{Å}, b = 11.24 \ \text{Å}, c = 14.13 \ \text{Å}$, $\alpha = 84.3^\circ$, $\beta = 112.6^\circ$, $\gamma = 134.6^\circ$, $q = 0.4a^*$, $V = 409.7 \ \text{Å}$.
| | | |

Fig. 5a – Isosurfaces of the 3D electrostatic potential map (e-map) extended along the stacking direction. Isosurface levels are $2\sigma[\Delta V(r)]$. b – [010]* e-map projection of the layer (1σ[ΔV(r)]) containing uranium, molybdenum and oxygen with its interpretation. The layer model is represented in the supercell $5a \times c$. c – Coordination polyhedra of U$^{6+}$ and Mo$^{6+}$. 

| a) | b) | c) |
|-----|-----|-----|
| **U-Mo-O** | **U-Mo-O** | **Ca-O** |
| **Ca-O** | **U-Mo-O** | **U-Mo-O** |
| **Ca-O** | **U-Mo-O** | **Mo** | **U** |

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were set according to the result of the chemical analysis in order to maintain the U:Mo:Ca ratio of 3:2:1. No modulation was applied to the atoms of the Ca–O layer as the refinement of this layer is not sensitive from our data. From the first refinement an insight of the calcium–water layer topology was revealed from the difference–Fourier map but it cannot be refined or described with accuracy. The final refinement leads to $R_{\text{obs}}/R_{\text{all}} = 0.3825/0.3834$ for 3953/17442 observed/all reflections (see Tab. 2). These values can be considered reasonable for the kinematical refinement of a modulated structure from electron-diffraction data (Boullay et al. 2013; Palatinus et al. 2015b; Steciuk et al. 2016). The refined structure is displayed on the Fig. 8. The results of the refinement are given in the Tab. 2 and the cation–oxygen distances, as a function of $t$, are presented in Fig. 9. The average bond valences for uranium are $BV(U1_1) = 6.146(5)$, $BV(U1_2) = 6.404(8)$, $BV(U2_1) = 6.196(7)$ and $BV(U2_2) = 6.011(6)$ vu.

5. Description of the structure

The structure of calcurmolite contains four symmetrically-distinct U sites, four Mo sites, two Ca sites and 24 O sites. In the structure of calcurmolite, all U is found to be hexavalent, occurring in pentagonal bipyramidal coordination with two strong U≡O bonds, (i.e. forming uranyl ion ($\text{UO}_2^{2-}$)) and five weaker bonds distributed in the equatorial plane (Figs 5b and 8; Tab. 3). The four symmetrically-distinct Mo sites are [6]-coordinated in the form of irregular, distorted octahedra (Fig. 5b and 8). The Mo1_1 and Mo1_2 (and Mo2_1 with Mo2_2) polyhedra share a common edge to form $\text{Mo}_2\phi_{10}$ dimers. The calcurmolite structure is based upon uranyl–molybdate sheets. The fundamental building blocks of those sheets are two trimers (I and II) of edge-sharing polyhedral composed respectively by two U1_1 and one U1_2 polyhedra (trimer I) and two U2_1 and one U2_2 polyhedra (trimer II) (Fig. 8). They form the six-membered clusters by sharing a common edge with other trimer (Fig. 8). These clusters are connected through the O5c (O5c_1 and O5c_2) equatorial atom of the U bipyramid to form irregular chains propagated along [–101]. These chains of polyhedra are connected through $\text{Mo}_2\phi_{10}$ dimers in a stair-case way. Sheets found in the structure of calcurmolite are of the same topology as observed in the structure of baumoite (Elliot et al. 2019).

Adjacent sheets are separated from each other at interplanar distance of 7.5 Å. Between the sheets are two symmetrically independent Ca atoms and four symmetrically independent H2O molecules (associated with Ow1_1, Ow1_2 and Ow2_1 and Ow2_2 atoms). The coordination of Ca atoms in the structure of calcurmolite is [6]. Most probably, Ow1_1 and Ow1_2 are transformer H2O group, with [3]-coordinated O atom, while Ow2_1 and Ow2_2 are non-transformer H2O group, with at least [4]-coordinated O atom (typology of H2O after Hawthorne and Schindler 2008).
The structure formula based on assumed occupancies is
$$\text{Ca}_{0.2}[(\text{UO}_2)_{0.6} (\text{MoO}_4)_{0.4} (\text{OH})_{0.8}] (\text{H}_2\text{O})_{~1.0}, \; Z = 4$$
in the modulated unit cell. In the supercell $5a \times b \times c$ giving a complete overview of the topology, the formula becomes
$$\text{Ca}[(\text{UO}_2)_{3} (\text{MoO}_4)_{2} (\text{OH})_{4}](\text{H}_2\text{O})_{~5.0}.$$ In his paper, Deliens (1992) defines calcurmolite more like a group of minerals having in common the presence of calcium, molybdenum, uranium and water. This assumption is explained by the discrepancies of the observed physical properties and variation in the powder X-ray diffraction patterns. The same diversity is found in the chemical formula reported by different studies (Rudnitskaya 1959; Fedorov 1963; Skvortsova et al. 1969; Deliens 1992; Sidorenko et al. 2005 and Dal Bo 2018). In the present work, the chemical formula is similar to the one reported by Dal Bo (2018) for calcurmolite from Rabejac Ca(UO$_2$)$_3$(MoO$_4$)$_2$(OH)$_4$(H$_2$O)$_8$. The difference in the water content can be due to the inaccuracy of the measurement in our study. Nevertheless, a variation in the amount of water or calcium seems highly possible considering the topology of the calcium–water layer and the disorder, especially along the stacking direction, observed in the diffraction data. It can probably occur without disturbing the topology of the uranium molybdenum layer and might be an explanation for the differences in the powder X-ray patterns reported in the literature.

6. Conclusions

The structure of calcurmolite can be considered complex regarding the topology and due to presence of commensurate/incommensurate modulations. The quality of the current data set does not allow differentiation between commensurate or incommensurate modulation but gives an insight of the structure for the first time. Although, the necessity of using the crenel-like (discontinuous) modulation functions indicates that the structure is close to be incommensurately modulated. Calcurmolite is similar to the Ba uranyl–molybdate baumoite, which was shown to be incommensurately modulated, having the same structural sheets, with U:Mo ratio equal to 3:2.
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Electronic supplementary material. Supplementary original crystallographic information file (cif) for calcurmolite is available online at the Journal web site (http://dx.doi.org/10.3190/jgeosci.297).

| atom       | harm. | x/a   | y/b   | z/c   | Uiso [Å²] | Delta/Occ. Delta | x4°  |
|------------|-------|-------|-------|-------|-----------|------------------|------|
| U1_1       | 0.5062| 0.9549| 0.3733|       | 0.0164(4) | 0.6              | 4.5  |
| Mo1_1      | –     | –     | –     | –     | –         | 0.0164(4)        | 4.9  |
| s, l       | 0     | 0     | –0.0158(5) |      |           |                  |      |
| c, l       | 0     | 0     | 0.0171(5)  |      |           |                  |      |
| U1_2       | –0.4966(16) | –0.9602(5) | –0.3912(3) |       | 0.0164(4) | 0.6              | 5.989(6) |
| Mo1_2      | –     | –     | –     | –     | –         | 0.0164(4)        | 4.989(9) |
| s, l       | 0     | 0     | –0.0087(4) |      |           |                  |      |
| c, l       | 0     | 0     | 0.0059(5)  |      |           |                  |      |
| U2_1       | 0.7517(19) | 0.9404(6)  | 0.0800(4)  |       | 0.0164(4) | 0.6              | 5.982(8) |
| Mo2_1      | –     | –     | –     | –     | –         | 0.0164(4)        | 1.0982(11) |
| s, l       | 0     | 0     | –0.0209(5) |      |           |                  |      |
| c, l       | 0     | 0     | –0.0296(5) |      |           |                  |      |
| U2_2       | –0.8693(17) | –0.9754(6) | –0.1270(3) |       | 0.0164(4) | 0.6              | 5.498(11) |
| Mo2_2      | –     | –     | –     | –     | –         | 0.0164(4)        | –0.280(2) |
| s, l       | 0     | 0     | 0.0078(4)  |      |           |                  |      |
| c, l       | 0     | 0     | 0.0022(5)  |      |           |                  |      |
| O5a_1      | 1.123(3) | 0.9597(5)  | 0.2324(5)  | 0.075(3) | 0.4       | 1.2467(16)       |      |
| O5a_2      | –0.948(4) | –0.93(1)  | –0.2544(5) | 0.075(3) | 0.4       | 0.018(2)         |      |
| O5b_1      | 0.330(3) | 0.9182(5)  | 0.2265(5)  | 0.075(3) | 0.4       | 0.9296(12)       |      |
| O5b_2      | –0.317(3) | –0.3938(5) | –0.2609(6) | 0.075(3) | 0.4       | 0.1708(15)       |      |
| O5c_1      | 0.602(3) | 0.8865(5)  | 0.2231(6)  | 0.075(3) | 0.4       | 0.4383(15)       |      |
| O5c_2      | –0.573(3) | –0.9107(5) | –0.2438(6) | 0.075(3) | 0.4       | 0.7681(17)       |      |
| O5d_1      | 0.515(3) | 0.9107(5)  | 0.2291(6)  | 0.075(3) | 0.2       | 0.7037(13)       |      |
| O5d_2      | –0.527(3) | –0.9321(5) | –0.2616(6) | 0.075(3) | 0.2       | 0.3868(16)       |      |
| O1a_1      | 0.006(2) | 0.2122(6)  | 0.374(2)   | 0.075(3) | 1        |               |      |
| O1a_2      | –0.032(1) | –0.2034(6) | –0.422(2)  | 0.075(3) | 1        |               |      |
| O1b_1      | –0.002(1) | 0.7229(7)  | 0.369(2)   | 0.075(3) | 1        |               |      |
| O1b_2      | 0.026(1) | –0.7229(6) | –0.369(2)  | 0.075(3) | 1        |               |      |
| O2a_1      | 0.266(1) | 1.1917(7)  | 0.123(2)   | 0.075(3) | 1        |               |      |
| O2a_2      | –0.402(1) | –1.2084(7) | –0.198(2)  | 0.075(3) | 1        |               |      |
| O2b_1      | 0.29(3)  | 0.6972(6)  | 0.056(2)   | 0.075(3) | 1        |               |      |
| O2b_2      | –0.343(1) | –0.7486(6) | –0.044(2)  | 0.075(3) | 1        |               |      |
| O3_1       | 0.2167(13) | 0.0220(3)  | 0.4328(5)  | 0.075(3) | 1        |               |      |
| O3_2       | –0.1301(18) | 0.0119(3)  | –0.4443(5) | 0.075(3) | 1        |               |      |
| O4_1       | 0.1117(17) | 0.8914(3)  | 0.0179(5)  | 0.075(3) | 1        |               |      |
| O4_2       | –0.293(2) | –1.0016(3) | –0.0730(4) | 0.075(3) | 1        |               |      |
| Cal_1      | 0.313(8)  | 0.351(2)   | 0.2608(14) | 0.104(1) | 0.4      |               |      |
| Cal_2      | –0.354(7) | –0.409(2)  | –0.3018(15) | 0.104(1) | 0.4      |               |      |
| Owl_1      | –0.608(10) | 0.480(3)   | 0.1181(19) | 0.104(1) | 1        |               |      |
| Owl_2      | 0.551(16) | –0.505(4)  | –0.1415(18) | 0.104(1) | 1        |               |      |
| Owl_2      | 0.192(9)  | 0.463(3)   | 0.3852(18) | 0.104(1) | 1        |               |      |
| Owl_2      | –0.136(8) | –0.479(3)  | –0.4143(16) | 0.104(1) | 1        |               |      |

Tab. 3 Positional parameters for calcurmolite (SSG P1(a000))
Fig. 9 The cations–oxygen distances (in Å) in the structure of calcurmolite.

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