Sequential dehydration of the phosphate–sulfate association from Gura Dobrogei Cave, Dobrogea, Romania

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Abstract. A rich association of primary guano minerals, including tatanakite, hydroxylapatite, brushite and gypsum with relics of illite, kaolinite, alpha (low) quartz and calcite, was identified in the fossil bat guano deposit from Gura Dobrogei Cave, Dobrogea County, Romania. Gypsum and Ca phosphates developed preferentially on the carbonate bedrock or on fallen carbonate blocks in the guano mass, whereas tatanakite was identified in the clay-rich, detritic sequences. The mineral species from the cave were characterized by optical methods, scanning electron microscopy, X-ray powder diffraction, Fourier-transform infrared and inductively coupled plasma–atomic emission spectrometry analysis. Chemically induced local dehydration of primary minerals, characterized by low temperatures (up to 100°C or even lower) and critically depending on exothermal reactions in the guano mass, prompted the formation of a secondary association, consisting of francoanellite, bassanite and monetite. Topotactic substitutions were observed in the cases of francoanellite on taranakite, basanite on gypsum and monetite on brushite. In its turn, ardealite was partially replaced by monetite and bassanite. The sequential dehydration process seems driven by the degradation of organic matter by microbial action and also, presumably, by other exothermic reactions at local scale (e.g., oxidation of ammonia, allogetic pyrite or other organic compounds).

1 Introduction

The sequential dehydration affecting guano masses in dry caves exposed to air circulation was described or supposed by a number of authors (e.g., Hill and Forti, 1997; Karkanas et al., 2000; Onac and Veřes, 2003; Frost and Palmer, 2011; McFarlane and Lundberg, 2018; Audra et al., 2019, and references therein) in relation to the thermal or dehydration reactions in the guano mass. Chemically induced mineral genesis is responsible for dehydration of previously formed mineral species and was considered a complementary tool for completing the archeological record (e.g., Karkanas et al., 2000). In spite of the utility of the mineralogical observation for understanding the cave behavior or for archeological records, very clear relations between the various “primary” and “secondary” mineral species were, however, rarely depicted in the same cave. Local dehydration of the very stable authigenic mineral tatanakite to francoanellite due to a local decrease in the water vapor pressure was, for example, described in the Măgurici Cave (Onac and Veřes, 2003) and expresses a path of the early diagenesis. In tropical caves where complete sequences of fossil bat guano minerals were subjected to desiccation, the microbial activity seems responsible of the breakdown by dehydration of variscite in berlinite or gibbsite (e.g., McFarlane and Lundberg, 2018).

Recent collecting in the fossil bat guano deposit in the cave from Gura Dobrogei (Romania) yielded several samples of great mineralogical interest. The aim of this work is to substantiate the data on the first described occurrence of monetite and the fifth occurrence of francoanellite in a Romanian cave and to offer new (mainly X-ray diffraction, XRD) data on a few more common cave minerals such as hydroxylapatite, brushite, ardealite, tatanakite, gypsum and bassanite (e.g., Hill and Forti, 1997; McFarlane and Lundberg, 2018; Audra et al., 2019). The mineralogical approach proposed in this paper intends to clarify the early diagenesis of the fos-
sil guano in one of the richest phosphate–sulfate associations from a Romanian Cave.

2 Geological setting

The cave from Gura Dobrogei is located in Constanța County, Dobrogea Region, Romania, about 3 km to the west from the homonymous locality and about 42 km north-northwest from Constanța, the major city in the area. The GPS coordinates of the main entrance (C in Fig. 1) are 44.467° N and 28.482° E, and its altitude is 106.7 m. The cave is also known as “the Bat Cave” (Peștera Liliecilor) from Gura Dobrogei.

The cave represents an upper (fossil) level of a karst system that developed on Visterna Brook, a tributary of Casimcea River, which in turn ends in Lake Tașaul, a former lagoon of the Black Sea. The cave has the main entrance on the right slope of the Visterna Brook and is developed in Upper Jurassic limestones of the Casimcea Syncline, in fact biostrome and bioherm limestones with sponges and microbialites (Bărăbulescu, 1999). The altitude of the main entrance is about 50 m from the thalweg of the brook. A detailed description of the cave and of its topography was given by Dumitrescu et al. (1958), from which was taken the plan view in Fig. 1.

The length of the main parts of the cave is about 480 m (Dumitrescu et al., 1958), but, after the investigation of the short and narrow divergent passages, the total length was 1365.8 m (https://www.speologie.org/pestera-liliecilor-de-la-gura-dobrogei, last access: 1 August 2020). A rich deposit of fossil bat guano is hosted in two of the galleries of the cave: the east–west oriented Gallery with Ceramics (or Circassian Gallery) and the north–south oriented Gallery with Fossils. The fossil guano deposit covers integrally the floor of the Gallery with Fossils, whereas the Circassian Gallery contains such a deposit only in its terminal part (Dumitrescu et al., 1958).

The humidity of the cave is about 80 %, and the mean temperature is 12.5 °C. The three entrances of the cave favor the formation of strong air currents that locally reduce the high humidity and dry the surface of the guano deposits. Some small colonies of insectivorous bats, pertaining to Mediterranean species Rhinolophus mehelyi and Myotis mystacinus, locally produce fresh guano accumulations. These recent accumulations increase the local temperature at the surface by
about 4 °C and also increase the content of moisture (28 to 33 vol %) and the acidity (pH = 5.19–5.96). In the fossil guano deposits, pH inversely correlates with depth and with moisture content. Moisture content measured for the fossil guano samples fluctuates between ~14 and 22 vol %, whereas pH values range from 6.18 to 7.15.

3 Analytical procedures

The samples were taken from both the Circassian Gallery and Gallery with Fossils, from both the surface and the mass of guano deposits, and from their contact with the wall rock and floor. The location of the sampling points is given in Fig. 1. In order to avoid supplementary dehydration, samples were stored after collection in sealed plastic bags at room temperature.

Observations of the crystal morphologies were conducted using a JEOL JSM-840 scanning electron microscope (SEM) manufactured by JEOL Ltd. (Tokyo, Japan). Acquisitions of energy-dispersive spectra (EDS) were made using a Tracer Northern TN-2000 system for 100 s (live time) with an accelerating voltage of 10 kV and a beam current of 10 nA. X-ray spectra were collected and processed with the PGT semiquantitative software. Full ZAF corrections (for atomic number, absorption and fluorescence effects) were applied to the raw X-ray data, but the results were of very poor quality due to the heating under the beam of the samples. The beam sensitivity due to dehydration prevented the acquisition of good-quality SEM pictures at high magnification.

Bulk methods of analysis were preferred, consequently. Samples were prepared for analysis by hand-picking bunches of crystals, and carefully verified for purity by X-ray powder diffraction. The purified separate was crushed, lightly ground of crystals, and carefully verified for purity by X-ray powder diffraction. The apparatus was operated under a voltage of 40 kV, with a beam current of 30 mA, using a step size of 0.04° 2θ, a counting time of 1 s per step and a scanning range of 5 to 90° 2θ. The slit system was the same for the other diffractometer used. Synthetic fluo- rite, with a = 5.4638(3) Å, was used as the internal standard. A complete set of data issued from up to 50 diffraction patterns is available upon request from the first author, whereas part of the X-ray powder data is given in the Supplement (Tables S1, S2). The unit cell parameters were refined from X-ray powder data by the least-squares program CELREF (Appleman and Evans, 1973) as modified for microcomputer use by Benoit (1987).

The infrared absorption spectrum of a francoanellite sample, the sole sample pure enough to be analyzed by this method, was recorded by a Fourier-transform Thermo Nicolet Nexus spectrometer (Thermo Fisher Scientific, Madison, United States), in the frequency range 400–4000 cm⁻¹, using a standard pressed-disk technique after embedding 1 mg of mechanically ground sample in 200 mg of dry KBr and compacting under 2500 N cm⁻² pressure.

Indices of refraction were measured without heating using calibrated oils, a spindle stage and a classical JENAPOL-U microscope (Carl Zeiss, Jena, Germany).

4 Mineralogy of the guano deposit

4.1 Primary (authigenic) minerals

4.1.1 Hydroxylapatite

Hydroxylapatite generally occurs as ochre to orange crusts lining the contact between the guano mass and the walls, as infillings of some veins traversing the guano mass, or as massive masses developed on the calcareous boulders contained by the guano. The crusts range up to 1 cm in thickness. The SEM examination shows that they are composed of thick beds of crystalline aggregates whose morphology varies from randomly deposited laths to post-colloidal, rosette-like de- posits. Intergrown laths with a hexagonal habit generally compose the aggregates. Individual crystals are platy on (0001) and typically smaller than 10 μm across; rarely they attain 20 μm. A typical aggregate of hydroxylapatite is shown in Fig. 2.

The indices of refraction for a representative sample (GDb 5 B), measured in immersion at 25 °C, are ω = 1.647(1) and ε = 1.640(1), which are identical to those given by Lehr et al. (1967) for the stoichiometric hydroxylapatite.

Unit cell parameters of hydroxylapatite are given in Table 1.
Two types of hydroxylapatite as follows: the crystallites in spite of a relatively good crystallinity. The hydroxyapatite from Gura Dobrogei. The arrow indicates a well-shaped crystal.

Table 1. Crystallographic parameters of selected guano minerals in the Gura Dobrogei Cave.

| Mineral species | Space group | Sample | \(a\) (Å) | \(b\) (Å) | \(c\) (Å) | \(\alpha\) (°) | \(\beta\) (°) | \(\gamma\) (°) | \(V\) (Å³) | \(n^1\) | \(n^2\) | \(2\theta^3\) |
|----------------|-------------|--------|--------|--------|--------|--------|--------|--------|--------|--------|--------|--------|
| Hydroxyapatite | \(P6_3/m\) | GDb 5 B | 9.427(3) | – | 6.856(4) | – | – | – | 527.7(4) | 7 | 38 | 10–80 |
| Hydroxyapatite | \(P6_3/m\) | GDb 7 A | 9.429(3) | – | 6.864(3) | – | – | – | 528.6(4) | 8 | 58 | 10–80 |
| Hydroxyapatite | \(P6_3/m\) | GDb 10 B | 9.428(4) | – | 6.877(5) | – | – | – | 529.3(4) | 6 | 42 | 10–80 |
| Brushite | \(Ia\) | GDb 3 A | 5.809(1) | 15.186(3) | 6.242(2) | – | 116.35(1) | – | 516.6(2) | 8 | 53 | 5–90 |
| Brushite | \(Ia\) | GDb 3 B | 5.815(2) | 15.181(5) | 6.241(2) | – | 116.29(2) | – | 504.5(3) | 9 | 62 | 5–90 |
| Brushite | \(Ia\) | GDb 5 A | 5.809(2) | 15.182(5) | 6.240(2) | – | 116.41(2) | – | 526.8(3) | 7 | 88 | 5–90 |
| Brushite | \(Ia\) | GDb 5 B | 5.840(3) | 15.192(7) | 6.252(3) | – | 116.49(3) | – | 542.1(3) | 5 | 35 | 5–90 |
| Brushite | \(Ia\) | GDb 7 A | 5.811(3) | 15.202(6) | 6.236(3) | – | 116.36(2) | – | 518.7(3) | 9 | 67 | 5–90 |
| Brushite | \(Ia\) | GDb 12 A | 5.840(4) | 15.204(8) | 6.259(3) | – | 116.29(3) | – | 508.9(4) | 5 | 55 | 5–90 |
| Brushite | \(Ia\) | GDb 13 A | 5.813(3) | 15.200(9) | 6.243(3) | – | 116.47(3) | – | 536.6(4) | 10 | 58 | 5–90 |
| Gypsum | \(I2/a\) | GDb 7 B | 5.717(3) | 15.198(4) | 6.517(1) | – | 118.27(2) | – | 498.8(3) | 6 | 43 | 10–85 |
| Gypsum | \(I2/a\) | GDb 12 B | 5.669(2) | 15.193(4) | 6.503(2) | – | 118.37(2) | – | 492.9(2) | 9 | 80 | 10–85 |
| Ardealite | \(Cc\) | GDb 3 A | 5.724(3) | 30.977(13) | 6.261(4) | – | 117.25(3) | – | 948.5(6) | 6 | 84 | 5–75 |
| Ardealite | \(Cc\) | GDb 5 A | 5.717(2) | 30.957(10) | 6.254(2) | – | 117.11(2) | – | 985.3(4) | 5 | 99 | 5–75 |
| Bassanite | \(I2\) | GDb 3 A | 12.046(9) | 6.921(8) | 12.749(9) | – | 90.25(6) | – | 1038.5(7) | 5 | 32 | 5–90 |
| Bassanite | \(I2\) | GDb 12 B | 12.099(3) | 6.919(1) | 12.689(4) | – | 90.46(2) | – | 1062.2(3) | 6 | 35 | 5–90 |
| Taranakite | \(R3c\) | GDb 6 A | 8.694(2) | – | 94.98(4) | – | – | – | 6218.2(313) | 7 | 92 | 5–66 |
| Francoanellite | \(R\bar{3}c\) | GDb 11 A | 8.695(2) | – | 82.44(2) | – | – | – | 5396.9(19) | 9 | 93 | 5–70 |
| Francoanellite | \(R\bar{3}c\) | GDb 11 A | 8.696(1) | – | 82.26(3) | – | – | – | 5388.1(18) | 6 | 62 | 5–90 |
| Monetite | \(P\bar{T}\) | GDb 3 A | 6.912(4) | 6.680(5) | 6.963(4) | 96.35(8) | 103.90(3) | 88.73(7) | 311.8(3) | 10 | 72 | 10–75 |
| Monetite | \(P\bar{T}\) | GDb 5 A | 6.680(5) | 6.667(9) | 6.919(7) | 94.17(3) | 104.06(6) | 87.25(8) | 307.3(8) | 7 | 98 | 10–75 |
| Monetite | \(P\bar{T}\) | GDb 13 A | 6.963(4) | 6.666(9) | 6.952(4) | 95.21(4) | 102.05(4) | 89.17(5) | 310.5(5) | 5 | 66 | 10–75 |

1 Number of cycles of refinement, 2 number of reflections used for refinement, 3 range of 2θ angles used in collecting the reflections.

Figure 2. SEM image showing a typical aggregate of hydroxyapatite from Gura Dobrogei. The arrow indicates a well-shaped crystal in the bottom-central part of the photograph. The scale bar (top) represents 10 µm.

Diffraction lines are slightly broadened by the fine size of the crystallites in spite of a relatively good crystallinity. The indices of crystallinity, calculated following the method proposed by Simpson (1964), differentiate, at Gura Dobrogei, two types of hydroxyapatite as follows:

– A first type, whose representative sample is GDb 5 B, is generally deposited directly on a carbonate support. The crystallinity index is high (I.C. = 0.324 for the referred sample), indicative of a poor crystallinity. Particularizing for the unit cell parameters in Table 1, they seem indicative of a carbonate-bearing hydroxyapatite since the parameter is smaller and the cell parameter is greater than those quoted by various authors for carbonate-free hydroxyapatite (e.g., \(a = 9.4176(5)\) Å and \(c = 6.8814(5)\) Å according to Elliott, 1994; \(a = 9.423(2)\) Å and \(c = 6.882(3)\) Å according to Bigi et al., 1996; \(a = 9.421(2)\) Å and \(c = 6.882(3)\) Å according to Brunet et al., 1999; \(a = 9.4244(2)\) Å and \(c = 6.8854(2)\) Å according to Morgan et al., 2000; and \(a = 9.4302(5)\) Å and \(c = 6.8911(2)\) Å according to Dorozhkin and Epple, 2002).

– A second type, whose representative samples are GDb 7 A and GDb 10 B, occurs in crusts differentiated in the guano mass. The crystallinity index is lower (I.C. = 0.077 and I.C. = 0.083, respectively, for the quoted samples), indicative of a high crystallinity. The unit cell parameters of the two samples of the second type maintain the characteristics of a carbonate-bearing hydroxyapatite.

4.1.2 Gypsum

Gypsum is one of the most common sulfates in the bat guano deposits from the caves (Hill and Forti, 1997). At Gura Dobrogei, the mineral occurs as centimeter-sized, white nodules or earthy masses in the guano layer and generally forms parallel aggregates of minute bladed crystals up to 100 µm in size.
length. The individual crystals are thin, flattened on \{010\} and elongated toward [001]. Stacking aggregates of crystals grown subparallel or parallel to \{010\} are common. The unit cell parameters determined for two representative samples by least-squares refinement, based on the best resolved X-ray powder diffraction lines, are given in Table 1. The refinements were carried out accepting the monoclinic symmetry, space group \(I2/c\), of the mineral (Cole and Lancucki, 1974).

### 4.1.3 Brushite

Brushite commonly occurs as very fine-grained snow-white coatings up to 1 cm thick covering hydroxylapatite, as irregular lining of hydroxylapatite bunches of crystals or as partial fillings of veinlets or cracks affecting the hydroxylapatite mass. The mineral also occurs as spheres up to 1 cm across enclosed by the detrital-rich sequences within the guano mass near calcite boulders. The SEM study shows that, in all cases, randomly oriented clusters of crystals with individuals reaching up to 15 µm in length (probably on [101] or [102] as for the synthetic crystals obtained by Abbona et al., 1993) compose the brushite aggregates. The most typical crystals are up to 10 µm long, 5 µm wide and 1 µm thick.

The unit cell parameters of brushite, calculated as the average of the seven sets of values in Table 1, are
\[
\begin{align*}
\alpha &= 5.820(14) \text{ Å}, \\
\beta &= 15.192(10) \text{ Å}, \\
\gamma &= 6.245(8) \text{ Å} \quad \text{and} \\
\beta &= 116.38(8)^\circ,
\end{align*}
\]
where the errors in the brackets correspond to the standard deviations of the mean (2σ) for each set of data. The obtained values are slightly larger than those given for the synthetic brushite by Curry and Jones (1971) (i.e., \(a = 5.812(2) \text{ Å}, b = 15.180(3) \text{ Å}, c = 6.239(2) \text{ Å} \quad \text{and} \quad \beta = 116.42(2)^\circ\)) but are in good agreement with those calculated by us for a sample of natural brushite from Angolo Incantato Cave, whose diffraction pattern was published by Balenzano et al. (1974): \(a = 5.818(1) \text{ Å}, b = 15.194(3) \text{ Å}, c = 6.247(2) \text{ Å} \quad \text{and} \quad \beta = 116.42(1)^\circ\).

### 4.1.4 Ardealite

Ardealite occurs as earthy, damp, white-yellow or off-white aggregates, directly deposited on hydroxylapatite crusts or as nests and small veins in the brushite–hydroxylapatite masses. These aggregates are composed of very fine, tabular sprays of crystals of maximum 20 µm in length and up to 1 µm wide. These compact radial groups are generally built up of thin blades or individual crystals with the \(c^*\) axis pointing outwards from the center. Stacking aggregates of crystals are also common.

As already mentioned by Dumitraș (2017), the X-ray powder diffraction pattern of ardealite is similar to, although distinguishable from, that of synthetic \(\text{Ca}_2\text{(HPO}_4\text{)}_2\text{(SO}_4\text{)}\cdot 4\text{H}_2\text{O}\) (Sakae et al., 1978). In all cases, at least four medium to strong lines in all the patterns, centered at about 4.30, 4.10, 3.15 and 2.95 Å, could not be indexed or related to any known impurity if we accept the crystallization of ardealite in the centrosymmetric group \(Cc\) (Sakae et al., 1978). The most representative X-ray diffraction pattern obtained for an ardealite sample from Gura Dobrogei was, however, indexed based on the \(Cc\) cell proposed by Sakae et al. (1978). The unit cell parameters (Table 1) are in good agreement with those reported by Sakae et al. (1978) for the synthetic \(\text{Ca}_2\text{(HPO}_4\text{)}_2\text{(SO}_4\text{)}\cdot 4\text{H}_2\text{O}\) \(a = 5.721(5), b = 30.992(5), c = 6.250(4) \text{ Å} \quad \text{and} \quad \beta = 117.26(6)^\circ\) and also with those refined for natural samples in the \(Cc\) hypothesis of symmetry by various authors (Dumitraș, 2017, and referred to works).

### 4.1.5 Taranakite

Taranakite generally forms chalky monomineralic nodular aggregates up to 1 cm across in the cave sediments resembling terra rossa. It also may form very fine powdery coatings on fractures affecting the sediment. In both cases taranakite masses are soft, earthy-looking and porous. The nodules are generally sharply defined because of the difference in color between them and the terra rossa mass. A transition zone where taranakite closely associates with α-quartz may locally be observed and occurs as coatings of powdery monomineralic taranakite nodules. Outside this zone, the unaffected sediment still contains illite and kaolinite. SEM microphotographs of taranakite aggregates (Fig. 3) show that they are composed of bunches of parallel slender laths or thin, roughly hexagonal platy crystals. Frequently, the crystals, which are only loosely interlocked within the masses, are poorly developed or broken. Individual crystals are flattened on \((0001)\), averaging 5 µm in size; they may exceptionally reach 15 µm across and 2 µm thick.

The unit cell parameters refined for a representative sample of taranakite from Gura Dobrogei are given in Table 1. The calculated \(a : c\) ratio (i.e., 0.0915 : 1), as well as the cell

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dimensions, fits well with the values reported by Marincea and Dumitraș (2003) for the synthetic potassium tarañakite [K₃Al₅(HPO₄)₆(PO₄)₁₂·18H₂O]. The unit cell parameters of tarañakite from Gura Dobrogei are smaller than the values refined for the synthetic K₃Al₅(HPO₄)₆(PO₄)₁₂·18H₂O \((a = 8.7025(11) \, \text{Å}, c = 95.05(1) \, \text{Å} \text{from X-ray powder data})\) but compare favorably with the values refined from neutron-diffraction data by Dick et al. (1998): \(a = 8.6882(3) \, \text{Å}, c = 94.98(2) \, \text{Å}\).

4.2 Secondary minerals resulting from dehydration

4.2.1 Bassanite

As well as in the cave from Cioclovina (Dumitraș et al., 2004), bassanite occurs as pseudomorphs after gypsum, whose perfect cleavage parallel to (010) is always observable. No textural signs of the reverse reaction, of bassanite hydration to gypsum (e.g., Van Driesche et al., 2012) were observed. The gypsum + bassanite association occurs as decimeter-sized, white nodules of earthy or chalky appearance included by the guano deposits. No fluorescence has been observed for these nodules under either short-wave (254 nm) or long-wave (366 nm) ultraviolet radiation. The SEM study shows that bassanite occurs as clustered acicular crystals that parallel the [001] axis of gypsum. They currently show parallel growth along the longest axis. Crystals are on the order of no more than 5 µm in length and often much less than this. Indices of refraction measured in yellow light \((\lambda = 589 \, \text{nm})\) as maximum and minimum values, respectively, by immersion of aggregates of platy crystals in calibrated liquids are \(n_{\min} = \alpha = 1.555(4)\) and \(n_{\max} = \gamma = 1.580(3)\). Accepting that the mean optical angle \((+2V_\alpha = 77.5^o)\) given by Palache et al. (1961) for the mineral is closely approximated in the case of our samples, it is possible to calculate a \(\beta\) value of 1.570, which is larger than that given for bassanite by Palache et al. (1961): \(\beta = 1.560\).

The unit cell parameters in Table 1 are in reasonable agreement with those refined for the stoichiometric calcium sulfate hydrate by Ballirano et al. (2001): \(a = 12.0350(5) \, \text{Å}, b = 6.9294(3) \, \text{Å}, c = 12.6705(4) \, \text{Å} \) and \(\beta = 90.266(3)^o\).

4.2.2 Francoanellite

In spite of the quite important number of francoanellite occurrences in bat guano deposits from Romanian caves (i.e., Onac and Vereş, 2003; Marincea et al., 2004; Giurgiu and Tămaș, 2013), there is a general lack of data on this rather rare mineral species. At Gura Dobrogei, francoanellite was identified as cream-white nodules of earthy appearance, clearly distinguishable in the terra rossa mass. In most of the samples, the mineral associates with tarañakite, illite 2M₁ and alpha quartz. The SEM study shows that francoanellite occurs in these cases as topotactic replacements of tarañakite (Fig. 3). The noticeable exception was encountered in the case of Sample GDb 11 A, in which all the diffraction reflections can be attributed to francoanellite. In both cases, the individual crystals are platy after (0001) and have hexagonal habit, with developments up to 10 µm across and 1 µm thick.

The indices of refraction were measured on stacks of platy subparallel crystals with hexagonal contour as maximum and minimum values, respectively. The average values obtained in yellow light \((\lambda = 589 \, \text{nm})\) are \(n_{\max} = \omega = 1.515(3)\) and \(n_{\min} = \epsilon = 1.509(2)\). The mean index of refraction that may be deduced from these values, applying the relation proposed by Mandarino (1976) for uniaxial crystalline compounds, is \(\bar{n} = (2\epsilon + \omega)/3 = 1.511\).

The average of five density measurements by the sink–float method, using a mixture of bromoform and toluene as immersion liquid, is \(D_m = 2.065(3) \, \text{g cm}^{-3}\). The calculated density for \(Z = 6\) (Dick and Zeiske, 1998), and considering a unit cell volume taken as the average of the two values in Table 1, is \(D_r = 2.268 \, \text{g cm}^{-3}\), which is practically identical to the measured value in the limit of errors. Note that all the measured values of density reported for francoanellite by previous authors (e.g., Balenzano et al., 1976) are slightly smaller than those calculated, which may be connected to the unusually high specific surface of the mineral and to its layered structure (Dick and Zeiske, 1998). Both values of density are smaller than that calculated for the synthetic K₃Al₅(HPO₄)₆(PO₄)₁₂·12H₂O by Dick and Zeiske (1998): \(D_r = 2.286 \, \text{g cm}^{-3}\), which essentially accounts for Na-for-K substitutions that prevail over the Fe³⁺-for-Al ones.

A bulk chemical analysis of francoanellite was done by ICP-AES using as starting material a practically monomineralic nodular mass (Sample GDb 11 A) surrounded by clay material. The analysis gave (in wt %) K₂O = 6.42, Na₂O = 3.19, MnO = 0.03, MgO = 0.05, CaO = 0.09, Al₂O₃ = 19.17, Fe₂O₃ = 2.11, P₂O₅ = 45.73 and H₂O (calculated) = 23.21 – total 100.00. All iron was taken as Fe³⁺. This composition, normalized on the basis of 22 cations in the anhydrous part of the compound and on 13 water molecules, leads to the following chemical-structural formula:

\[
[H_6.000K_{1.690}Na_{1.275}Ca_{0.020}Mg_{0.015}Mn_{0.005}]Al_{4.658}Fe_{0.327}(PO_4)_{8.000}·13H_2O.
\]

Compared with the type material from Castellana Cave, Italy (Balenzano et al., 1976), francoanellite from Gura Dobrogei is richer in both sodium and aluminum, which corroborates the presence of kaolinite and the local abundance of colloidal iron sesquioxides and Na-rich clay material (montmorillonite?) in the terra rossa mass. The structural position normally occupied by K⁺ ions is slightly overcompensated for with a cation sum of 3.005 vs. the ideal 3.00, which indicates that the (NH₄)⁺-for-K substitution is absent or very limited.

The chemical refractive energy (\(K_c\)) value is 0.2279 and is based on the formula given before and the Gladstone–Dale constants (Mandarino, 1981). The physical refractive energy (\(K_p\)) value, as derived from the average refractive index (\(\bar{n}\))
Figure 4. Infrared spectrum of francoanellite from Gura Dobrogei Cave.

and the calculated density is 0.2253. The physical refractive energy calculated on the basis of the measured density is $K_p' = 0.2256$. The compatibility index $(1 - K_p/K_c)$ is 0.011 (for $K_p$) and 0.010 (for $K_p'$), which is, in both cases, rated as superior (Mandarino, 1981).

Representative diffraction patterns for the purest francoanellite samples identified at Gura Dobrogei were collected with both Cu $K\alpha$ and Fe $K\alpha$ radiation. They are available under request from the first author, and a representative XRD pattern is given in Sect. S1 in the Supplement. The unit cell parameters refined for the two samples (GDb 11 A and GDb 11 B, respectively) are given in Table 1. They are slightly smaller than the values for $H_6K_3Al_5(PO_4)_8\cdot13H_2O$ ($a = 8.71$ Å, $c = 82.50$ Å according to Smith and Brown, 1959) which could account for limited Fe$^{3+}$-for-Al replacements. On the other hand, both sets of values compare favorably with the value refined on the basis of X-ray data for the same compound by Dick and Zeiske (1998): $a = 8.6897(16)$ Å and $c = 82.271(13)$ Å. The unit cell parameters refined by us are also close to those that can be refined on the basis of the data published for francoanellite from the type locality (Grotte di Castellana, Italy), i.e., $a = 8.697(7)$ Å and $c = 82.43(2)$ Å (as refined on the basis of the two sets of XRD values given by Balenzano et al., 1976), but are slightly smaller than those of francoanellite from Grotta della Rondinella (Italy): $a = 8.721(3)$ Å and $c = 82.91(6)$ Å (as determined on the basis of the three sets of XRD values given by Balenzano et al., 1979).

Figure 4 gives an infrared spectrum of a sample of francoanellite from Gura Dobrogei, whereas Table 2 lists the wavenumbers, characters and intensities of the bands, together with their assignments.

A few comments regarding the spectrum can be made as follows. (1) There is a quite strong resemblance between the infrared spectrum of francoanellite and that of taranakite, based on structural parallelism between the two mineral species (Dick et al., 1998; Dick and Zeiske, 1998). (2) As well as in taranakite (Marincea and Dumitras, 2003), in the O-H stretching region, the infrared absorption spectrum shows one broad and two narrow absorption bands, whose frequency and width indicate hydrogen-bonded O-H stretching vibrations, corresponding to two different positions of the molecular water. In its turn, the H-O-H “scissors” bending of H$_2$O is double degenerated (Table 2). (3) No additional vibration band which may be assigned to the $\nu_4$ in-plane bending vibration mode of the (NH$_4$)$^+$ structural group occurs at $\sim 1430$ cm$^{-1}$, indicating the absence of the (NH$_4$)$^+$-for-K$^+$ substitutions. (4) As in taranakite, the tetrahedral phosphate anions (part of them protonated) occupy two crystallographic non-equivalent positions, but no signs of polymerization of two protonated phosphate groups according to schema: $2(HPO_4)^{2-} = (H_2PO_4)^- + (PO_4)^{3-}$ can be observed, which agrees with the structure resolved by Dick and Zeiske (1998). (5) The antisymmetric stretching of the unprotonated phosphate groups is not double degenerated like in taranakite (Ross, 1974; Marincea and Dumitras, 2003).
Table 2. Positions and assumptions of the infrared absorption bands recorded for francoanellite from Gura Dobrogei.

| Structural group | Vibrational mode                      | Wavenumber (cm⁻¹) | Intensity, character¹ |
|------------------|---------------------------------------|-------------------|-----------------------|
| (HPO₄)²⁻         | O-H stretching                        | 3453              | s, sh                 |
| H₂O              | O-H stretching                        | −                  | b                     |
| (HPO₄)²⁻         | O-H stretching                        | 2960              | m, b                  |
| H₂O              | H-O-H bending²                        | 2462              | m, b                  |
| H₂O              | H-O-H bending²                        | 1679              | m, sh                 |
| (HPO₄)²⁻         | δ O-H in-plane bending²               | 1657              | m, sh                 |
| (HPO₄)²⁻         | ν₃ O-P-O antisymmetric stretching²    | 1203              | s, sh                 |
| (HPO₄)²⁻         | ν₃ O-P-O antisymmetric stretching²    | 1118              | s, sh                 |
| (PO₄)³⁻          | ν₁ O-P-O antisymmetric stretching     | 1073              | s, sh                 |
| (PO₄)³⁻          | ν₁ O-P-O symmetric stretching         | 1008              | s, sh                 |
| (PO₄)³⁻          | P-O-H symmetric stretching            | 985               | s, sh                 |
| (PO₄)³⁻          | ν₁ O-P-O symmetric stretching²       | 934               | s, sh                 |
| (HPO₄)²⁻         | P-O-H symmetric stretching            | 844               | m, shd                |
| (HPO₄)²⁻         | δ O-H out-of-plane bending            | 840               | m, sh                 |
| (HPO₄)²⁻         | δ O-H out-of-plane bending            | 775               | w, sh                 |
| (HPO₄)²⁻         | ν₄ O-P-O in-plane bending             | 697               | w, b                  |
| (HPO₄)²⁻         | ν₄ O-P-O in-plane bending             | 649               | m, sh                 |
| (PO₄)³⁻          | ν₄ O-P-O in-plane bending             | 554               | s, sh                 |
| (PO₄)³⁻          | ν₄ O-P-O in-plane bending             | 516               | m, shd                |
| (AlO₆)⁹⁻ (?⁻)    | O-VIAl-O stretching                   | 461               | s, sh                 |
| (HPO₄)²⁻         | ν₂ O-P-O out-of-plane bending         | 455               | s, shd                |
| (PO₄)³⁻          | ν₂ O-P-O out-of-plane bending         | 416               | m, sh                 |
| (HPO₄)²⁻         | ν₂ O-P-O out-of-plane bending         |                    |                       |
| (HPO₄)²⁻         | ν₂ O-P-O out-of-plane bending         |                    |                       |

¹ s = strong; m = medium; w = weak; vs = very strong; sh = sharp; b = broad; shd = shoulder; ² assumptions coincident with those 
given for tanzanite by Arlidge et al. (1963) and Ross (1974); ³ may also represent a H₂O libration or a lattice mode.

### 4.2.3 Monetite

Monetite was identified in samples containing brushite, and rarely ardealite, as a dehydration product of brushite. Macroscopically, the mineral can not be distinguished properly from the admixed brushite, but its presence was always confirmed by XRD study. The preservation of the samples in sealed plastic bags at room temperature excludes an accidental dehydration. The mixed brushite–monetite samples preserve the snow white macroscopic color of brushite. The SEM study of a monetite-rich aggregate (Sample GDb 5 A) showed groups of platy individual crystals elongated on an indefinite direction of up to 8 µm in length and up to 3 µm large. Because of the high volatility of the sample under the electron beam, it was impossible to obtain fair-quality microphotographs. It is, however, obvious that the monetite aggregates do not substantially differ from the brushite ones, suggesting that the topo tactic substitution of brushite by monetite, following the brushite dehydration, was the principal mechanism of monetite formation.

The crystals of monetite are too small to properly ascertain the refraction indices. For this reason, these were measured by immersion in Cargille oils, as maximum and minimum values, on composite, carefully hand-picked bundles of crystals that show a brushite-like tabular habit. The values obtained were \( n_{\text{max}} = 1.64 \) and \( n_{\text{min}} = 1.59 \), which closely approach those for \( \gamma = (1.640–1.65) \) and \( \alpha = (1.587–1.60) \), respectively (Palache et al., 1961).

The unit cell parameters refined for three representative samples of monetite on the basis of reflections in the XRD pattern unequivocally attributable to this mineral are given in Table 1, whereas a representative pattern is given in Sect. S1. The structure resolved by Catti et al. (1980) was accepted for the unit cell parameters given in Table 1. The structure resolved by MacLennan and Beever (1955) and a = 6.90(1) Å, \( b = 6.65(1) \) Å, \( c = 7.00(1) \) Å, \( \alpha = 96.35^\circ \), \( \beta = 103.90^\circ \) and \( \gamma = 88.73^\circ \) according to MacLennan and Beever (1955) and \( \alpha = 6.910(1) \) Å, \( b = 6.627(2) \) Å, \( c = 6.998(2) \) Å, \( \alpha = 96.34(2)^\circ \), \( \beta = 103.82(2)^\circ \) and \( \gamma = 88.33(2)^\circ \) according to Dickens et al. (1971), respectively. The cell orientation with \( c > a > b \) established by refinement (MacLennan and Beevers, 1955; Dickens et al., 1971) is not respected by two but one of the samples in Table 1.

### 4.3 Relict minerals in the guano mass

Quartz, calcite, 2M1 illite and 2M kaolinite were identified as relict minerals in the guano mass. Their description, as well as the crystallographic parameters refined on the basis of XRD patterns, is given in the Supplement (Sect. S2).
All the “primary” phosphates and sulfates (i.e., hydroxyapatite, brushite, ardealite, tikanakite and gypsum) are clearly authigenic, resulting from normal interactions between the pre-existing sediments from the cave floor and walls and the acidic solutions derived from guano. In fact, the biologically driven reactions from the guano mass, and particularly the bacteria-induced ones, ultimately release phosphoric and sulfuric acids (e.g., Forti, 2001; Onac and Forti, 2011). Additional generation of sulfuric acid may have occurred from pyrite or hydrogen sulfide oxidation sourced from green schist formation at the base of the Upper Jurassic limestones contained in the cave. The strongly acidic character of the solutions favors the crystallization of both primary phosphates and sulfates and of the monoclinic polytype of kaolinite (Fialips et al., 2000). The presence of (SO$_4$)$_2$ in solution is clearly critical for the formation of guano and ardealite.

Takanakite may be interpreted to have been derived through the in situ fixation of potassium and aluminum ions into a phosphate structure (e.g., Sakae and Sudo, 1975). As well as the aluminum, the alkali ions may be available rather from local clay minerals that are susceptible to partial dissolution by the low-pH solution developed from the guano deposit than from distally sourced groundwater. The relics of illite and kaolinite in the tikanakite plus francoanellite mass, as well as the presence of a low crystallinity, semi-colloidal alpha quartz, suggest the decomposition of clay minerals by acidic solutions.

Brushite was probably formed both through replacement of hydroxyapatite and directly by reaction of the acidic solutions derived from the bat guano with calcium carbonate though mechanisms similar to those proposed by Frost and Palmer (2011) or Pak (1981) (see below). Bassanite, francoanellite and monetite formed during a later stage of guano evolution, typical for the “dry” karst systems, and were found in the driest parts of the cave.

Presumably organic matter from the bat guano depositions, and particularly ammonia, gave locally strong exothermic reactions capable of raising the temperature to up to 100°C. In fact the reaction of oxidation of ammonia is strongly exothermic (being characterized by a strong negative enthalpy), making it an important local heat source. This reaction can be ideally written as follows:

$$4\text{NH}_3(g) + 5\text{O}_2(g) \rightarrow 4\text{NO}(g) + 6\text{H}_2\text{O}(g) (\Delta H = -905.2 \text{kJ mol}^{-1})$$

The principal source of ammonia is the urea in the bat droppings. Urea eliminates easier ammonia than hydrolyzes, resulting in quite high local concentrations of NH$_3$ (e.g., Alexandrova and Jorgensen, 2007). Ammonia can also result from reactions similar to that proposed by Frost and Palmer (2011) or Pak (1981) for the formation of brushite from an ammonia-bearing precursor, e.g.,

$$(\text{NH}_4)_2\text{HPO}_4 + \text{CaCO}_3 + \text{H}_2\text{O} \rightarrow \text{CaHPO}_4 \cdot 2\text{H}_2\text{O} + \text{CO}_2 + 2\text{NH}_3.$$
tallization of bassanite plus monetite of up to 100 °C is more than reasonable.

6 Conclusions

Based on the textural relations between the various mineral species, the result is that the chemical and then the thermal transformation of the guano mass at local scale during an early stage of diagenesis was responsible for the structure of the mineral associations in the Gura Dobrogei Cave. The mineral distribution and the local textures suggest that the advance of the reaction front affecting the guano mass was controlled by the general porosity of the sedimentary matrix. The acid solutions that percolated the guano mass may be residual products of microbial degradation that are mostly acids (e.g., Alexander, 1977). The dehydration process of primary guano minerals seems driven by the degradation of organic matter by microbial action and also, presumably, by other exothermic reactions at local scale (e.g., oxidation of ammonia, allogenic pyrite or other organic compounds). In all cases, the breakdown of the organic material is the driving force for authigenic mineral formation.

Data availability. The paper uses GIR data. GIR does not allow for redistribution except for the purpose of replication archives. Permissions can be obtained upon request to the second author.

Supplement. The following materials are available in the Supplement: Sect. S1: representative X-ray powder data for secondary minerals from Gura Dobrogei Cave; Table S1: X-ray powder data for a selected sample of francoanellite from Gura Dobrogei Cave; Table S2: X-ray powder data for a selected sample of monetite from Gura Dobrogei Cave; and Sect. S2: retic minerals in the guano mass (Sect. 4.3.) The supplement related to this article is available online at: https://doi.org/10.5194/ejm-33-329-2021-supplement.

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