ON SOME CAVE MINERALS FROM NORTHERN NORWAY

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ABSTRACT

The present paper aims to point out the results of 31 samples from some Norwegian caves that have been analysed with respect to their mineralogical composition. Identification of the minerals were done by X-ray diffraction, thermal, infrared spectroscopy and scanning electron microscopy. Seventeen minerals were identified belonging to 4 groups: carbonates, sulphates, oxides-hydroxides and silicates.

INTRODUCTION

The most representative karst phenomenona in Norway are to be found around and above the Arctic Circle, being developed in metamorphic marbles and dolomites (Lauritzen, 1991). The karst landscape from north Norway offer an overwhelming variety of morphological features on all scales, which have resulted in a wide range of publications within geomorphology, hydrology, sedimentology and chronology.

Cave minerals have not previously been described in detail. The cold conditions of Norwegian caves associated with a variety of minerals interbedded in layers with the marble strata, and an extremely diverse petrography of the rocks which come in contact with the marbles could provide favorable conditions for the formations of a large variety of cave minerals. The present paper presents the analysis of 31 mineral samples from 15 locations.

Due to the cold conditions and high latitude, speleothems are rare in Norwegian caves when compared with caves from temperate zones. This is in part due to climatic (temperature, Pco₂) controls (Ford & Williams, 1992), and to lithofacial factors (Onac, 1991).

The name and location of the sampled caves are withheld for conservational reasons.

MINERALOGICAL ASPECTS

Thirty-one samples from different Norwegian caves had been analysed so far in respect to their mineralogical composition. X-ray diffraction (XRD), as well as thermal, infrared spectroscopy (IR) and scanning electron microscopy (SEM) were used for this purpose. The analysis were done in two different laboratories belonging to the University of Bergen (Norway) and University of Cluj (Romania) using Philips and JEOL devices assisted by computers.

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Normally, three main categories of minerals are found in caves: (1) **allogenic minerals**, transported into the cave by mechanical action of gravity, water or wind, (2) **authogenic residual minerals** (fragments of bedrock, minerals formed during petrogenesis and debris left behind from the dissolution processes) and (3) **authogenic precipitates** formed *in situ* by chemical processes in the cave system. Our paper deals mostly with the latter one.

1. **Carbonates**

**Calcite** is by far the most common carbonate mineral in the Norwegian caves, being the stable form of calcium carbonate under the temperature, total pressure and CO$_2$ partial pressure of the cave environment. Aragonite and hydromagnesite can be found as well.

Calcite frequently appears to occur as a single mineral which exhibit almost infinitely varied morphologies. The most representative are flowstone, dripstone, rimstone, pool deposits and moonmilk. All of these form in almost any cave.

Moonmilk is a very common speleothem in many Norwegian caves (Onac, 1991). It is a soft, white (sometimes white-yellowish) mixture of microscopic crystals of different carbonates or sulphates with an important amount of water (Hill & Forti, 1986). The moonmilk samples we analysed were mostly calcitic and in just one case we found hydromagnesite and gypsum. As shown by SEM (Fig. 1) the calcite crystals appear to form either fibrous-lamellar or acicular-prismatic microcrystals.

In a previous paper, Onac & Farcas (1992), pointed out large moonmilk deposits located along well ventilated galleries, which are composed entirely of thread-shaped crystals (fibrous-lamellar or acicular prismatic) (Bernasconi, 1975) in size between 8 and 50 cm.
80 µm in length and 0.1-2 µm in thickness. A primary origin (Diaconu, 1976) was ascribed to these deposits which have been precipitated directly from super-saturated solutions. The moonmilk which has been sampled from dry cave passages consist of paramorphs of calcite after aragonite and more often a consolidation of moonmilk

Fig. 2 - SEM. Moonmilk in which calcite crystals suffered a cimentation process (sample 810).

Fig. 3 - SEM. Lamellar aggregates of hydromagnesite (sample 764).
aggregates by new forming calcite is seen (Fig. 2). The cave ventilation has a major influence on the dynamics of the precipitation processes by setting up local thermodynamic conditions. The morphological differences we noticed among calcite crystals composing our sampled moonmilk reflect these various thermodynamic conditions (Onac & Ghergari, 1993).

In a moonmilk-like sample collected from behind some “gypsum balloons” we found *hydromagnesite*. It forms very thin scaly crystals arranged in the shape of flakes (Fig. 3), showing symmetric extinction and positive elongation (Onac & Ghergari, 1993). Mirror-image twins on (100) with elongate lamellae along the c-axis have also been observed.

White, powdery deposits (on the cave floor) of hydromagnesite were also observed in one cave. We believe that the occurrence of hydromagnesite can be explained either by seepage of magnesium-rich solutions into the cave, when soft moonmilk precipitate or by magnesium waters seeping up through cave soil with deposition of white earthy-like deposits.

The mineralogical investigations carried out on some light-gray, well crystallized crusts, have pointed out microaggregates of acicular crystals which were analysed by means of XRD technique. The spectrum (Fig. 4) indicated the presence of *aragonite*. Closely associated with aragonite, we found tabular gypsum crystals.

Aragonite can nucleate and grow quite easily when the necessary supersaturation
Fig. 6 - SEM. Gypsum moonmilk; lamellar crystals (sample 765).

Fig. 7 - SEM. Tabular crystals of thenardite (sample 766).
level is attained and when the growth of calcite is suppressed. For this it seems that the magnesium, strontium and sulfate ions must poison the crystal growth of calcite, to allow the supersaturation level where aragonite can precipitate (White, 1976). Of these ions, sulfate (in gypsum) and magnesium (from wall-rock) are present in the close vicinity and can be the prime factor influencing aragonite deposition in preference to calcite deposition.

2. Sulphates

Among sulphate minerals, gypsum is very common in many Norwegian caves, as our analysis pointed out. Three forms are present: (1) gypsum crusts, (2) euhedral crystals grown in a clay matrix and (3) gypsum moonmilk.

Both granular and fibrous crust varieties have been found: the latter often display thin-walled semispherical balloon-like crust, several centimeters across. Behind these balloons a white crystalline powder was collected. The XRD spectrum (Fig. 5) indicated a mixture of celestite and gypsum crystals. To confirm the identification of celestite we have made some observation at the polarizing microscope. Transparent to translucent celestite grains, with vitreous to pearly luster having strong relief (when compared with gypsum) were observed. It seems that celestite has been precipitated by percolating waters carrying strontium in solution leached from the wall rocks.

Euhedral gypsum crystals take a columnar shape (001), sparsely prismatic, usually perfectly transparent. Sometimes they are slightly coloured by fine grains of clay. Needle-like and sword crystals are to be found penetrating the clay layer’s surface. Their size do not exceed 1 cm in length and 0.2-1 cm in thickness.

The gypsum moonmilk consists of lamellar microcrystals with silky lustre that form delicate aggregates having different habits and sizes (0.1-1.5 mm). Closely associated with gypsum we found large grains with dull lustre ascribed to calcite. The crystals’
The morphology analysed by SEM (Fig. 6) showed that gypsum is a neomorphic mineral having euhedral habit (grains displaying fully developed crystal form) while the calcite appears as relict clasts (grains that present corroded edges).

The other sulphate mineral we found is thenardite; this mineral is stable in a dry cave environment (for more details see Hill & Forti, 1986, p.114). Thenardite was first time seen as white microcrystalline aggregates spread over a thick porous floor soil deposit that covers one room, often used by speleologists as a camp site. SEM analysis of this microcrystalline material showed an aggregate consisting of tabular crystals, developed parallel to (001) with more or less corroded edges (Fig. 7). The EDX spectrum irrefutably proved the presence of this mineral (Fig. 8).

Due to the high level of human impact (camping and cooking), in this place the humidity is well under the normal values which characterize the cave environment (over 90%) being possible for this reason a direct precipitation of thenardite. In most of the caves where this mineral was found (except lava tubes), originally, precipitated as mirabilite and dehydrating to thenardite when the cave environment became drier.

We consider oxidation of sulphides (i.e. pyrite and marcasite, especially) locally within the cavernous limestone, as well as, oxidation of sulphides from elsewhere in the stratigraphic section and transport into the cave by percolating water to be the main sources of $\mathrm{SO}_4^{2-}$. No exhaustive investigations have been done with respect to the presence of thenardite, but the following mechanism can be possible: an interaction between the cave floor sediments with high content of muscovite and orthoclase which could provide $\mathrm{Na}^+$ and sulphate solutions moving upwards through that porous cave soil.

4. Oxides and Hydroxides

In a cave environment, pH is generally between 7 and 8 (with few exceptions), Eh is in the range of +0.4 to +0.6 volts (oxidizing) and activity of water is near unity (Hill & Forti, 1986). With respect to this situation, the stable form of iron is a ferric hydrate and for manganese a mixed $\mathrm{Mn}^{3+}$ and $\mathrm{Mn}^{4+}$ oxide. Because these compounds are highly insoluble they tend to appear as crusts and coatings but generally do not form speleothems themselves.

We sampled from different caves dull opaque, brownish-black to black crusts looking very similar to what we know to be called wad. Our analysis revealed the same composition as in most of the references (e.g. Kashima, 1983). “Wad” is typically a fine-grained mixture, chiefly pyrolusite and romanechite, both being identified through

![Fig. 9 -XRD spectrum of todorokite from sample 808a.](image)
XRD analysis in our samples as well. Another complex hydrous manganese oxide was identified by laboratory study (chemical, XRD and IR) to be todorokite (Fig. 9).

Their presence in caves can be ascribed to the weathering process of manganeseous carbonates or silicates which are common in the surrounding area of the sampled caves. Bacteria may also play a part in their genesis (Coman, 1984).

Reddish-brown to brownish-yellow coatings and small rimstone dams made up of goethite are to be found in many caves covering either calcite or sand speleothems. Most of the common yellow-brown ferric oxides usually known as limonite properly belong to this species. Limonite is usually associated with acidic groundwaters. Limonite and goethite speleothems occur in caves where the cavernous limestone underlies iron-bearing rocks or where rich iron rock lenses are to be found interbedded in marble. Oxidation of siderite and pyrite-rich horizons produces acidic solutions which can retain the iron in solution. When these solutions are neutralized after they have percolated downward into the cavernous limestones, the iron is precipitated. Limonite and goethite can be transported in suspension as colloids or as very fine-grained particles until the material accumulates in dripstone deposits. All minerals ascribed to this group, mentioned until now, formed under oxidizing conditions.

The crystalline forms of solid water (ice) belong as well to the oxide minerals. Given the high latitude of the Norwegian karst, and because of the mean annual temperature that in some places falls below 0°C, both ephemeral ice, forming in the entrance passages of caves during the cold seasons and perennial ice occurring when special airflow conditions maintain low temperatures during the warm season, are to be found.

The variety of ephemeral ice speleothems is very rich, although not as rich as that of calcite speleothems. They resemble one another in many ways, especially where the forms are the direct result of flowing water (stalactites, stalagmites, ice falls).

Caves containing perennial ice are known as glacieres. The ice stratigraphy could provide extremely important paleoenvironmental informations by using 14C to date organic matter lenses (wood, bone, etc.) that are interbedded in the ice. Measuring d18O of the formation water it is possible to determine the absolute temperature changes that occurs over a time interval.

5. Silicates

There is not much to say about silicate minerals in the Norwegian caves as long as they are mainly clays, usually transported as clastic sediments and hardly correspond to the definition of speleothems and cave minerals that always formed after the development of the cavities. We identified in our samples: quartz, muscovite, orthoclase, tremolite, kaolinite and palygorskite. The presence of these minerals is due either to weathering processes of mica schist and of marble (i.e. authigenic weathering residues) or because they were carried unchanged into the cave from outside (alloogenic residues).

We found them associated with manganese oxides and calcite speleothems as well. Any of these mentioned silicate minerals do not form speleothems themselves.

CONCLUSIONS

Mineralogically, the Norwegian caves we investigated contain 17 different minerals so far, assigned to four chemical groups. Although the best represented group is that one of oxides-hydroxides, the minerals found in it form very few speleothems (an exception
being the ice and goethite) when compared with the great variety of speleothems formed by the minerals from the carbonates and sulphates group.

The extensive calcitic moonmilk deposits found in many caves from north of Norway prove that the underground microclimate (very cold and wet) is favourable to the deposition of this speleothem. The outlook of the microcrystals present in both calcitic and gypsum moonmilk highly depends on the local thermodynamic conditions.

The sequence of allogenic and autogenic residual minerals is well represented (6 species). Their importance arise from being a basic material which could provide, through its weathering, ions that can further react and form new cave minerals.

Because the human impact on the cave environment almost always leads to irreversible destruction and just sometimes (very rare) to useful changes (see our thenardite), we have to protect the underground world in which minerals, animals and fossils are extremely well preserved.

The present investigation was intended as a first identification and description of the cave minerals forming speleothems in few caves from northern Norway. Further work on this topic is required to answer questions of origin, depositional mechanisms and the way the cave environment has to be protected against any kind of outside influence, so that the cave microclimate will rest undisturbed.

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REFERENCES


