Ghiaraite: A new mineral from Vesuvius volcano, Naples (Italy)

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ABSTRACT

In this work we report the first finding of CaCl₂β·4H₂O, long known as a synthetic phase. The mineral, called ghiaraite, was discovered in 2011 in a sample belonging to the Real Museo Mineralogico di Napoli (Italy), that had been collected in 1872 at Vesuvius volcano and stored in a glass sealed vial. It is associated with chlorocalcite (KCaCl₃), hematite, sylvite, and halite. The mineral was found inside an ejecta of 5 m in size transported by a lava flow to the locality of Massa di Somma. Here with the ejecta still hot the sample was collected and rapidly stored in a sealed glass vial to preserve it from the atmospheric conditions. Ghiaraite is triclinic, space group *P*Ī, with unit-cell parameters: a = 6.3660(5), b = 6.5914(5), c = 8.5568(6) Å, $\alpha = 93.504(6)^\circ$, $\beta = 97.778(7)^\circ$, $\gamma = 110.557(6)^\circ$, V = 330.802(9) Å³, Z = 2. The calculated density is 1.838 g/cm³ using the ideal formula and the powder X-ray diffraction data. It occurs as euhedral isometric grains up to 5–6 µm long intimately intermixed with chlorocalcite. The eight strongest reflections in the X-ray powder diffraction pattern [listed as d(Å)(I)(hkl)] are: 2.628(100)(022); 2.717(88)(103); 4.600(88)(111); 2.939(77)(200); 2.204(75)(121), 5.874(73)(100), 6.124(47)(010); 3.569(46)(111).

Ghiaraite was approved by the Commission on New Minerals, Nomenclature and Classification with IMA number 2012-072. The mineral was named in honor of Maria Rosaria Ghiara (b. 1948), Head of Real Museo Mineralogico of Napoli and Centro Musei delle Scienze Naturali e Fisiche dell'Università degli Studi di Napoli Federico II for her important work in promoting the scientific research focused on the mineralogy of Vesuvius volcano.

Keywords: Ghiaraite, new mineral, X-ray diffraction, EDS, Vesuvius volcano, calcium tetrahydrate chloride

INTRODUCTION

The new mineral described in this work, named ghiaraite, appears to be extremely rare and was found by one of the authors (M.R.) in association with chlorocalcite (KCaCl₃), hematite, sylvite, and halite in the mineral collection "Vesuviana" of the Real Museo Mineralogico di Napoli in 2011. Ghiaraite occurs in voids in the ejecta surface that were exposed to volcanic vapors. The sample was collected at the Vesuvius volcano, stored in a sealed glass vial and deposited at the Real Museo di Napoli in 1872 by Arcangelo Scacchi (1810-1894), a member of the "Reale Accademia delle Scienze Fisiche e Matematiche," see Figure 1. More in detail, it was found inside an ejecta of 5 m in size transported by a lava flow to the locality of Massa di Somma (coordinates: 40.5° N, 14.22 E). Here, with the ejecta still hot (Scacchi 1874), the sample was collected and within a few minutes stored in a sealed glass vial to preserve it from the atmospheric conditions. Arcangelo Scacchi reported the

description of the ejecta finding on a note published on October 1872 right after the Vesuvius eruption of April 26, 1872 (Scacchi 1872). The note was published by the "Società Reale di Napoli" in the "Rendiconto dell'Accademia delle Scienze Fisiche e Matematiche" (year XII). In the note of 1872 Scacchi described chlorocalcite with the wrong chemical formula CaCl; however, he reported a chemical analysis in which the CaCl compound represented the 58.76% with the remaining 32.24% constituted by potassium chloride, sodium chlorides, and manganese chlo-



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FIGURE 1. Original sealed glass vial deposited by Scacchi in 1872 at Real Museo di Napoli. The vial is about 5 cm long.

ride. At that time the microprobe analysis was not an available technique and the data reported by Scacchi were clearly a bulk chemical analysis on a mixed sample. Therefore, Scacchi could not identify ghiaraite by its chemical composition, and could not realize that "chlorocalcite" was actually KCaCl₃ (Zambonini 1935). Chlorocalcite is also obtained a synthetic compound by melting a 1:1 mixture (molar ratios) of KCl and CaCl₂ at 750 °C (Korshunov et al. 1984). Similarly, the synthetic analog of ghiaraite was first reported in 1875 to be obtained while melting CaCl₂·6H₂O (Gmelin 1957).

In this work we report the first natural finding of CaCl₂·4H₂O, which has been named ghiaraite in honor of Maria Rosaria Ghiara (b. 1948) who is Head of Real Museo Mineralogico of Napoli and Centro Musei delle Scienze Naturali e Fisiche dell'Università degli Studi di Napoli Federico II. Several years ago Ghiara began a program to promote scientific collaborations to obtain new information about the stored minerals from the Vesuvius volcano in the last 200 years.

Ghiaraite was approved by the Commission on New Minerals, Nomenclature and Classification with IMA number 2012-072. The type specimen is deposited in the collections of the Real Museo of Napoli with registration number 16986-E5525.

APPEARANCE, MORPHOLOGY, PHYSICAL AND OPTICAL PROPERTIES

Ghiaraite occurs as euhedral isometric very small grains (up to 5–6 μ m long) intimately intermixed with chlorocalcite (Figs. 2 and 3). The bulk sample chlorocalcite–ghiaraite appears milky white toward light gray in color. The forms cannot be easily determined as ghiaraite appears as intergrown crystals. The association chlorocalcite–ghiaraite is strongly deliquescent when exposed to the air.

The combination of small grain size plus sensitivity to the moisture did not allow us to obtain a better physical or optical characterization, with the exception of the calculated density, which is 1.838 g/cm³ using the ideal formula and the unit-cell volume obtained from powder X-ray diffraction data. For the synthetic α -CaCl₂·4H₂O analog, the measured densities are 1.826(3) g/cm³ at 25 °C (Wulff and Schaller 1934), and 1.837 g/cm³ (Thewalt and Bugg 1973). These data are in perfect agreement with the calculated density of our sample.

From the data obtained on the synthetic α -CaCl₂·4H₂O analog, Bunn et al. (1935) deduced that the compound is biaxial negative, not pleochroic, and with refractive indexes $\alpha = 1.530$, $\beta = 1.557$, $\gamma = 1.567^{\circ}$. The only available data, even if of poor quality, are those from Wulff and Schaller (1934) who reported n_{min} = 1.54(8), n_{max} = 1.56(6). No optical properties were measured on ghiaraite as the typical crystal size did not allow any optical experimental measurement.

EXPERIMENTAL METHODS

X-ray powder diffraction

X-ray powder diffraction (XRPD) data were recorded on a PANalytical θ - θ diffractometer (Cu radiation) equipped with a long fine-focus Cu X-ray tube operating at 40 kV and 40 mA and a Real Time Multi-STRIP (RTMS) detector (X'Celerator). The active length (detector aperture) of the X'Celerator (RTMS detector) is 2.122° and this leads to collect the signal over a large angular range at the same time, reducing the total time of the scan. The data collection time is about



FIGURE 2. SEM image of the intermixed sample of chlorocalcite, ghiaraite, sylvite, and hematite (large single crystal at the bottom of the sample).

100 times shorter with respect to an identical scan performed using the traditional proportional detector (Reiss 2002). The data were collected in the angular range $3-67^{\circ}$ in 2θ in continuous mode with an integrated step scan of 0.033° in 2θ and a total time for the data acquisition of about 8 min (angular speed $0.14^{\circ}/s$). This corresponds to a scan with a virtual counting statistic of 30 s/step. The program High Score Plus (PANalytical) was used for phase identification and Rietveld refinement (Rietveld 1967). The structural model of synthetic CaCl₂·4H₂O (ICSD code 8061, PDF 01-070-2220), determined by Leclaire and Borel (1979) was used as starting structural model of ghiaraite. A pseudo-Voigt function was employed for the profile shapes. Refined parameters (Gaussian and Lorentzian coefficients).

SEM and EDS

The SEM-EDS investigation was carried out using a CamScan MX2500 scanning electron microscope, equipped with a LaB₆ cathode, four quadrant solid-state BSE detector and a EDAX EDS system for micro-analysis. The analytical conditions were: accelerating voltage of 20 kV, filament emission of ~130 μ A, and working distance of 27 mm.

RESULTS AND DISCUSSION

Ghiaraite was found during an X-ray diffraction investigation in collaboration between the Real Museo Mineralogico di Napoli, the Department of Geosciences of University of Padova, and the Museo of Mineralogy of University of Padova, with the aim to identify the rare minerals collected at the end of 1800 at Vesuvius volcano. In detail, the aim of this collaboration was initially that to identify the chlorocalcite phase. The original glass vial (Fig. 1) containing ghiaraite and chlorocalcite was cut and a small fraction of the sample was collected extremely quickly to avoid the deliquescence phenomenon occurring for such phases. X-ray diffraction data were measured very quickly thanks to the use of the highly sensitive detector available and by SEM-EDS under vacuum. The results of the study are reported below.

X-ray diffraction

The sample analyzed by X-ray diffraction is constituted by chlorocalcite, ghiaraite, and sylvite (Fig. 4) (the fourth associated phase is hematite, but this mineral was found inside the vial as separate crystals not intermixed with chlorocalcite, ghiaraite, and sylvite). Patterns of synthetic α -CaCl₂·4H₂O strongly agreed with



FIGURE 3. Enlargement of Figure 2 with evident ghiaraite (isometric and euhedral grains) and chlorocalcite (surrounding matrix).

the peaks of natural ghiaraite. Ghiaraite is triclinic with space group $P\overline{1}$. The Rietveld refinement using the structural data of the synthetic compound gives the following results: the agreement indices were $R_p = 5.09\%$ and $R_{wp} = 6.68\%$. For ghiaraite, $R_{Bragg} =$ 1.22%. Data of *d*-spacing (in Å for CuKa) are provided in Table 1, whereas the refined unit-cell parameters are the following: *a* = 6.3660(5), *b* = 6.5914(5), *c* = 8.5568(6) Å, *a* = 93.504(6), $\beta =$ 97.778(7), $\gamma = 110.557(6)^{\circ}$, *V* = 330.802(9) Å³, *Z* = 2. Leclaire and Borel (1979) report for their synthetic sample the following cell parameters: *a* = 6.3673(8), *b* = 6.5932(6), *c* = 8.5606(15) Å, *a* = 93.50(1), $\beta = 97.83(1)$, $\gamma = 110.58(1)^{\circ}$. Thewalt and Bugg (1973) report: *a* = 6.364(5), *b* = 6.593(2), *c* = 8.557(3) Å, *a* = 93.52(4), $\beta = 97.77(5)$, $\gamma = 110.56(3)^{\circ}$. Such excellent match

Chemistry

Due to the impossibility of preparing a suitable sample for WDS analyses, we could only qualitatively analyze the sample of Figures 2 and 3 by SEM-EDS. The EDS spectra are shown in Figure 5. It can be noted that for ghiaraite we nearly could not measure the water oxygen as the vacuum under which the measurements were performed is in the order of the 10⁻⁵ mbar. Under such vacuum conditions, the coordinated water of a compound like CaCl₂·4H₂O is totally lost. In addition, the limited amount of potassium visible in the spectra for ghiaraite must belong to chlorocalcite, this is quite evident by the X-ray diffraction data: in fact the unit-cell volume of ghiaraite only differs from the synthetic calcium-chloride tetrahydrated by less than 0.1%; this is in agreement with the total absence of a large cation like K⁺, which would strongly affect the unit-cell volume. Consequently, the combined study of ghiaraite by EDS and X-ray diffraction definitively prove that ghiaraite is the natural analog of the tetrahydrated calcium chloride, CaCl₂·4H₂O, for which the crystal structure has been well known for about 40 years (i.e., Thewalt and Bugg 1973).

Relation to other species

Ghiaraite has the synthetic analog α -CaCl₂·4H₂O, which can be obtained during the melting of CaCl₂·6H₂O as well as by crystallization from aqueous solutions. However, it can be formed by simple hydration of anhydrous CaCl₂ or CaCl₂·2H₂O. It is also formed by interconversion of other polymorphs (β - and γ -CaCl₂·4H₂O, both showing *P*2₁/*c* space group), as it appears to

FIGURE 4. Diffractogram of the sample containing ghiaraite (indicated with symbol G) (reference diffractogram: ICSD card no. 8061). In green and in brown chlorocalcite (ICSD card no. 00-021-1170) and sylvite (ICSD card no. 98-009-9788) are shown.

	mewaite	ina bagg	(1)))		
	Ghiaraite			Thewalt and Bugg (1973)	
hkl	$d_{\rm obs}$	$I_{\rm rel}$	$d_{\rm calc}$	$d_{\rm calc}$	I _{rel}
010	6.124	47	6.130	6.130	53
100	5.874	73	5.875	5.874	99
1 1 0	5.315	29	5.317	5.315	31
111	4.600	88	4.603	4.599	100
111	3.569	46	3.568	3.568	56
0 1 2	3.296	38	3.295	3.295	42
1 2 1	2.998	44	2.996	2.996	40
200	2.939	77	2.938	2.937	64
103	2.717	88	2.719	2.716	78
022	2.628	100	2.629	2.629	84
2 2 1	2.574	29	2.573	2.571	21
2 2 1	2.499	20	2.499	2.499	16
121	2.396	36	2.397	2.397	33
022	2.350	25	2.350	2.350	20
222	2.300	24	2.301	2.300	19
121	2.204	75	2.204	2.204	76
1 3 1	2.090	33	2.090	2.090	31
014	2.067	23	2.067	2.067	20

TABLE 1. X-ray powder-diffraction data for ghiaraite including the

 d-spacings calculated using the crystal structure data by

 Thewalt and Bugg (1973)

be the most stable tetrahydrated form (Gmelin 1957). It is itself a hygroscopic compound and its occurrence and role in the system CaCl₂–H₂O has been thoroughly studied by Basset et al. (1933, 1937) and Lannung (1936).

The analog of ghiaraite was found by Chesnokov in a burning coal dump at mine no.45 in Kopeisk, Chelyabinsk Oblast, Southern Ural, Russia, and described as unnamed "tetrahydrated calcium chloride" (Chesnokov et al. 1987) and then under the name "mesohydrite" (Chesnokov et al. 1994). "Mesohydrite" was described as a product of dehydration of antarcticite; physical properties and X-ray powder data identical to synthetic CaCl₂·4H₂O were reported (Chesnokov et al. 1994). However, the above locality has a clearly technogenic character and "mesohydrite", thus, was considered as a substance of anthropogenic origin and never approved as a valid mineral species by the IMA. On the contrary ghiaraite described in this work has undoubtedly a natural origin.

The structure of the synthetic analog of ghiaraite by Thewalt and Bugg (1973) shows Ca^{2+} be coordinated by $3Cl^- + 4H_2O$, forming $Ca_2Cl_4(H_2O)_8$ molecular moieties that are held together by only weak H-bonds and van der Waals forces. This is quite different from the structures of antarcticite ($CaCl_2 \cdot 6H_2O$) (Torii and Ossaka 1965), in which columns of face-sharing $Ca(H_2O)_9$ polyhedra are linked through Cl^- ions, or sinjarite (Aljbouri and Aldabbagh 1980), in which $CaCl_4(H_2O)_2$ octahedra share corners to form $CaCl_2(H_2O)_2$ sheets, linked via H-bonds. Ghiaraite can be considered an intermediate hydrate of $CaCl_2$, between sinjarite and antarcticite. There does not appear to be any isostructural minerals.

Sonnenfeld and Kühn (1993) noted the presence of a mineral with this formula in an evaporite from Sergipe, Brazil, and introduced the name sergipite for it. But no mineralogical data were presented.

The natural origin of ghiaraite: Monitoring the CaCl₂ and chlorocalcite (KCaCl₃) kinetics

To definitively demonstrate that ghiaraite has a natural origin we investigated the kinetic behavior of a synthetic CaCl₂ crystal-

FIGURE 5. Qualitative analysis by EDS of chlorocalcite and ghiaraite performed on the sample shown in Figure 2.

line compound under the same temperature and humidity conditions usually present in laboratory (23 °C and 30% of relative humidity). About 150 mg of powder CaCl₂ were deposited above a zero-background sample holder (in pure silicon) and kept for five hours at 300 °C to have an anhydrous starting material. The temperature and the relative humidity were monitored observing fluctuations of about 1 °C and 1% of relative humidity.

Very fast X-ray powder diffraction analyses were carried out, with each diffractogram collected over 3 min between 10 and 46° in 20. The measurements were performed in a continuous mode over five hours. Considering the "zero minute" the exact instant when the sample was extracted from the furnace, the first X-ray measurement was performed after 3 min: after only 3 min the sample resulted to be constituted by about 70% of CaCl₂ and 30% of CaCl₂·2H₂O (the quantitative analyses were performed by method). The natural analog of CaCl₂·2H₂O is called sinjarite (Aljbouri and Aldabbagh 1980). The diffractograms showing the kinetics investigated are plotted in Figure 6a: it is possible to note that CaCl₂ (indicated with symbol A) decreases gradually to disappear after 36 min, completely transforming to sinjarite (indicated with symbol B). After 39 min, sinjarite starts to transform to ghiaraite $[CaCl_2 \cdot 4H_2O]$, indicated with symbol T in Figure 6b. After 1 h and 3 min, sinjarite and ghiaraite are present at 50%, and only after 1 h and 51 min sinjarite completely disappears (Fig. 6c). From 1 h and 51 min on, only ghiaraite remains but it starts to become amorphous.

The second step of our kinetic analysis was focused on the

FIGURE 6. X-ray powder diffractograms relative to the kinetics of anhydrous CaCl₂ (symbol A), sinjarite (symbol B), and ghiaraite (symbol T).

Position [°2Theta] (Copper (Cu))

25

30

20

possibility that ghiaraite could be originated by the chlorocalcite decomposition after the extraction of the sample from the ejecta by Scacchi in 1872. Therefore, we synthesized a sample of chlorocalcite (prepared by melting a 1:1 mixture of KCl and CaCl₂ at 750 °C for 2 h). As done for the above experiments, the X-ray measurements were performed very quickly with single measurements of 3 min each. After 1 h and 5 min chlorocalcite was totally decomposed to KCl and an amorphous compound and sinjarite and/or ghiaraite never crystallized.

Concerning ghiaraite, we should recall (see experimental method section at X-ray powder diffraction) that after the opening of the vial in which the original sample was kept and the preparation of the X-ray experiment the total time necessary to obtain the diffractogram was 8 min. Vial opening, sample preparation, and X-ray analysis were performed under the same identical conditions described above (23 °C and 30% of relative humidity). Our kinetics studies clearly indicate that ghiaraite in 8 min could not be formed by hydration of CaCl₂ or sinjarite.

Scacchi described that he stored the ghiaraite-bearing sample extracted from the ejecta very quickly (a few minutes or even less) and even if a small amount of air remained trapped in the vial it was not sufficient to hydrate sinjarite to from ghiaraite as chlorocalcite, the strongly dominant phase, acted as a very strong buffer as demonstrated by our kinetic experiments.

ACKNOWLEDGMENTS

The research was supported by "Progetto d'Ateneo 2006, Università di Padova" to F.N.

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MANUSCRIPT RECEIVED MAY 13, 2013 MANUSCRIPT ACCEPTED SEPTEMBER 1, 2013 MANUSCRIPT HANDLED BY FERNANDO COLOMBO