## **Article**

# Napoliite, Pb<sub>2</sub>OFCl, a new mineral from Vesuvius volcano, and its relationship with dimorphous rumseyite

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#### **Abstract**

Napoliite, ideally Pb2OFCl, is a new fluoroxychloride mineral found in specimen from a



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fumarole formed subsequent to the 1944 eruption of Vesuvius volcano, Naples Province, Italy. It occurs as well-shaped lamellar crystals up to  $0.25 \times 0.25 \times 0.01$  mm typically forming clusters up to  $0.4 \times 0.4$  mm on the surface of volcanic scoria in association with anglesite, artroeite, atacamite, calcioaravaipaite, cerussite, challacolloite, cotunnite, hephaistosite, manuelarossiite, matlockite, and susannite. Napoliite is colourless with white streak and adamantine lustre. It is brittle and has a laminated fracture. Cleavage is perfect on  $\{001\}$ .  $D_{\text{calc}} = 7.797$  g cm<sup>-3</sup>. The calculated mean refractive index is 2.10. Chemical composition (wt.%, electron microprobe) is: PbO 91.71, F 3.89, Cl 7.34, -O=(F+Cl) -3.30, total 99.64. The empirical formula calculated on the basis of 3 anions is Pb<sub>1,999</sub>O<sub>0,997</sub>F<sub>0,996</sub>Cl<sub>1,007</sub>. Raman spectroscopy confirms the absence of OH<sup>-</sup> groups and H<sub>2</sub>O molecules in the mineral. Napoliite is tetragonal, space group  $P4_2/mcm$ , a =5.7418(11), c = 12.524(4) Å, V = 412.9(2) Å<sup>3</sup> and Z = 4. The strongest lines of the powder X-ray diffraction pattern [d, Å (I, %) (hkl)] are: 3.860 (85) (111); 3.139 (20) (004); 2.914 (100) (113); 2.866 (63) (200); 2.118 (19) (204); 2.027 (19) (220); 1.665 (20) (313); 1.642 (23) (117). The crystal structure was refined to  $R_1 = 0.024$  for 222 reflections with  $F > 4\sigma(F)$ . It is based on lead oxide blocks derived from that of litharge PbO which alternate with layers of chloride ions. Napoliite represents a new structure type with a unique order/disorder pattern of fluorine and oxygen atoms. The new mineral is dimorphous with rumseyite. It is named after the city of Naples (Napoli in Italian).

**Keywords**: napoliite; new mineral; lead oxyhalides; crystal structure; rumseyite; fumarole sublimate; Vesuvius volcano.

## Introduction

The Vesuvius volcano, located about 9 km east of Naples in Campania, Italy is one of the world's most famous geological objects and attracts millions of visitors each year. It is also one of the most studied volcanoes and is a world-class mineralogical location regardless. To date, the Somma-Vesuvius volcanic complex is the type locality of 68 mineral species. While most of them have "grandfathered" status, several minerals have been described more recently as a result of new finds or the re-examination of old samples facilitated by progress in X-ray diffraction and other analytical techniques. Among them are d'ansite-(Mn), Na<sub>21</sub>Mn<sup>2+</sup>(SO<sub>4</sub>)<sub>10</sub>Cl<sub>3</sub> (Demartin *et al.*, 2012), ghiaraite, CaCl<sub>2</sub>·4H<sub>2</sub>O (Rossi *et al.*, 2014), parascandolaite, KMgF<sub>3</sub> (Demartin *et al.*, 2014), tondiite, Cu<sub>3</sub>Mg(OH)<sub>6</sub>Cl<sub>2</sub> (Malcherek *et al.*, 2014), verneite, Na<sub>2</sub>Ca<sub>3</sub>Al<sub>2</sub>F<sub>14</sub> (Balić-Žunić *et al.*, 2018), sbacchiite, Ca<sub>2</sub>AlF<sub>7</sub> (Campostrini *et al.*, 2019), and paradimorphite, As<sub>4</sub>S<sub>3</sub> (Campostrini *et al.*, 2022). Herein, we report another new mineral, napoliite named after the city of Naples (Napoli in Italian).

The new mineral, its name and symbol (Npi) have been approved by the Commission on New Minerals, Nomenclature and Classification of the International Mineralogical Association (IMA2022-073; Kasatkin *et al.*, 2023). The holotype specimen is deposited in the collections of the Fersman Mineralogical Museum of the Russian Academy of Sciences, Moscow, Russia with the registration number 5885/1. A part of the holotype specimen is deposited in the collections of the Sezione di Mineralogia "Alessandro Guastoni" of the Museo della Natura e dell'Uomo, University of Padova, under catalogue number MMP M23222.

## Occurrence and mineral association

The Somma-Vesuvius complex (Campania region, Italy) is characterized by complex magmatism and volcanism (Lima *et al.*, 2007; Alfano and Parascaldola, 2015; Sbrana *et al.*, 2020; Melluso *et al.*, 2022); it is a polygenic stratovolcano with strong eruption variability in terms of style, cyclicity and magma composition. During the last 3.5 ka effusive lava flows and scoria eruptions were alternated to highly explosive Plinian eruptions of pumice and ash, with pyroclastic flows and surges. All exposed lavas refer to the recent activity between a.d. 1631 and 1944 (Santacroce *et al.*, 2008; Sbrana *et al.*, 2020). Since 1944, the Vesuvius has been quiescent, with only moderate seismicity and fumarolic activity.

The last eruption dating back to 1944, began on March 18<sup>th</sup> and ended on the 29<sup>th</sup> of the same month, marked the transition from an open conduit to a closed conduit state. The eruption took place during wartime that precluded the collecting of specimens after the eruption and their investigation. Antonio Parascandola was the only person who studied the mineralogy of these fumaroles between 1948 and 1960, when temperatures decreased from a maximum of 800°C in 1950 to about 460°C in 1960s (Parascandola, 1951, 1960, 1961). His studies revealed the presence of about ten mineral species new for Vesuvius.

In recent times, the systematic study of fumarolic minerals formed after the 1944 eruption was undertaken by a joint team of scientists from the Department of Chemistry of the University of Milan and the Naples section of the National Institute of Geophysics and Volcanology – Vesuvius Observatory (see, *e.g.*, Campostrini and Gramaccioli, 2005; Russo and Campostrini, 2011; Russo *et al.*, 2014) that resulted in a discovery of several rare minerals early unknown at this locality, such as ammineite, artroeite, caledonite, fluornatrocoulsellite, gearksutite, hemimorphite, and matlockite and the description of two new species – parascandolaite, KMgF<sub>3</sub> (Demartin *et al.*, 2014) and sbacchiite, Ca<sub>2</sub>AlF<sub>7</sub> (Campostrini *et al.*, 2019).

Napoliite, therefore, is the third new mineral discovered in the material from fumaroles of the 1944 eruption. The sample of lava scoria where napoliite occurs was collected by one of the authors (MR) in early 2010s from a large fragment located at the eastern rim of the "Gran

Cono" crater. Napoliite overgrows volcanic scoria and is associated with anglesite, artroeite, atacamite, calcioaravaipaite, cerussite, challacolloite, cotunnite, hephaistosite, matlockite, susannite, and the recently discovered new mineral manuelarossiite PbCaAlF<sub>7</sub> (IMA2022-097, Nestola *et al.*, 2023). Hephaistosite and susannite are of special interest, since both of them were found on Vesuvius for the first time. Hephaistosite occurs as small grains up to 0.02 mm forming a limited solid solution series with Tl-bearing challacolloite. The most Tl-rich hephaistosite variety contains (wt.%): K 2.04, Tl 16.06, Pb 57.47, Cl 24.08, total 99.65. Its empirical formula is  $(Tl_{0.58}K_{0.38})_{\Sigma 0.96}Pb_{2.04}Cl_{5.00}$ . Susannite occurs as prismatic colourless crystals up to 1mm long with adamantine lustre. Its identification is based on semi-quantitative chemical analysis (only Pb, S, C, O detected at EDS with Pb to S ratio equal to 4:1) and single-crystal X-ray diffraction data. The parameters of its trigonal unit cell are: a = 9.080(10), c = 11.565(12) Å, and v = 826.7(15) Å<sup>3</sup>.

## General appearance and physical properties

Napoliite occurs as well-shaped lamellar crystals up to  $0.25 \times 0.25 \times 0.01$  mm typically forming clusters up to  $0.4 \times 0.4$  mm on the surface of volcanic scoria (Fig. 1). The new mineral is colourless and transparent with white streak and adamantine lustre. It is brittle and has a laminated fracture. No fluorescence is observed under long- or short-wave ultraviolet light. The hardness is  $\sim 3$  on the Mohs' scale. Cleavage is perfect on  $\{001\}$ . The density of napoliite could not be determined due to the scarcity of available material. Density calculated using the empirical formula is  $7.797 \text{ g cm}^{-3}$ .

Napoliite is optically uniaxial. Its refractive indices (RI) could not be measured because of the absence of immersion liquids that can measure RI values higher than 2.0. The Gladstone-Dale relationship predicts an average RI of 2.10 which is similar to that of other Pb oxychloride minerals [2.13 for blixite Pb<sub>8</sub>O<sub>5</sub>(OH)<sub>2</sub>Cl<sub>4</sub> (Gabrielson *et al.*, 1958; Krivovichev and Burns, 2006); 2.27 for yeomanite Pb<sub>2</sub>O(OH)Cl (Turner *et al.*, 2015); 2.32 for damaraite Pb<sub>3</sub>O<sub>2</sub>(OH)Cl (Criddle *et al.*, 1990), *etc.*]. Optical properties of napoliite were studied using the methods common for metallic minerals. In reflected light, napoliite is grey with no visible bireflectance and pleochroism. In crossed polars it is very weakly anisotropic. Internal reflections are not observed. The set of reflectance measurements performed in air relative to a SiC standard by means of a Universal Microspectrophotometer UMSP 50 (Opton-Zeiss, Germany) is given in Tab. 1 and plotted in Fig. 2 in comparison with the published data for rumseyite (Turner *et al.*, 2012).

#### Raman spectroscopy

The Raman spectrum of napoliite (Fig. 3) was collected in the range 50–400 cm<sup>-1</sup> as no peaks were observed above 350 cm<sup>-1</sup>. The data were collected using a DXR Thermo Scientific Raman spectrometer, equipped with a diode-pumped solid state 532 nm laser. The analytical points were performed with a 50× LWD (long working distance) objective, operating at a power of 4 mW, with a spectral resolution in the range of about 3 cm<sup>-1</sup> and a spatial resolution of about 1 μm. The acquisition time adopted was 10 seconds for 10 scans accumulation.

The Raman spectrum of napoliite shows two very intense peaks at 144 and 130 cm<sup>-1</sup>, whereas lower intensity peaks are positioned in the region between about 315 and 330 cm<sup>-1</sup>, and even lower intensity peaks at about 180, 112, 83, 60 and 56 cm<sup>-1</sup>. No Raman signals are detected for those regions above 350 cm<sup>-1</sup> including the OH<sup>-</sup> region above 3000 cm<sup>-1</sup>.

The Raman spectra of synthetic compounds structurally related to napoliite are unknown. In addition, the Raman spectrum of its polymorph rumseyite is also unreported. However, we can compare the napoliite Raman spectrum with the F-free synthetic compound Pb<sub>3</sub>O<sub>2</sub>Cl<sub>2</sub> described by Zakir'yanov *et al.* (2016). Based on this work, the peaks in the 130 –150 cm<sup>-1</sup> region of napoliite could be assigned to the Pb–Cl stretching vibrations and the peaks in the 315 – 330 cm<sup>-1</sup> region to the Pb–O stretching vibrations. However, the strongest peak in the Raman spectrum of litharge, along with another strong peak at 339 cm<sup>-1</sup>, is observed at 146 cm<sup>-1</sup> (Ciomartan et al., 1996). Thus, the peak at 144 cm<sup>-1</sup> may be a superposition of the bands of Pb–Cl stretching and O–Pb–O bending vibrations.

On the other hand, an alternative interpretation of the Raman spectrum of napoliite in the range  $110 - 150 \text{ cm}^{-1}$  is possible. The vibrational mode, which corresponds to the most intense band with the wavenumber  $k_{\rm w} = 144 \text{ cm}^{-1}$ , belongs to the ground vibrational state (zeroth vibrational level). At room temperature, the corresponding normal vibrations are partially excited. The fractions of Pb–O bonds located at the first and second vibrational levels (without taking into account anharmonicity) are  $p_1 = \exp[-(hv)/(kT)]$  and  $p_2 = \exp[-(2hv)/(kT)]$  of the fraction of the unexcited state, respectively, where h is the Planck constant,  $v = ck_{\rm w}$  is the frequency (c is the speed of light in vacuum), k is the Boltzmann constant and T is temperature. According to the relationship  $hv = kT_{\rm eq}$ , the temperature  $T_{\rm eq}$  of 298 K is equivalent to the wavenumber  $kw^{298}$  of 207 cm<sup>-1</sup>, and the values of  $p_1$  and  $p_2$  are equal to 0.5 and 0.25, respectively. Considering that the polarizability (and hence the intensity of Raman scattering) depends on the anharmonicity, it can be assumed that the bands at 139 and 112 cm<sup>-1</sup> refer to thermally excited states at the first and second vibrational levels with anharmonic shifts of 14 and 32 cm<sup>-1</sup>, respectively.

Based on data from Zakir'yanov *et al.* (2016), the peaks at 112 and 180 cm<sup>-1</sup> are assigned to the O–Pb–O bending vibrations and the peaks at 56, 60 and 83 cm<sup>-1</sup> to Cl–Pb–O, Cl–Pb–F

and Cl–Pb–F bending vibrations. Comparison with Raman spectra of oxygen-free lead halides are generally in agreement with the assignment made by Zakir'yanov *et al.* (2016). Raman spectra of all lead chlorides and fluorides do not contain bands of fundamental modes above 240 cm<sup>-1</sup> whereas such bands are observed in Raman spectra of the Pb<sup>2+</sup> oxides, litharge (at 321 and 339 cm<sup>-1</sup>: Ciomartan *et al.*, 1996; Madsen and Weaver, 1998) and massicot (at 289 and 384 cm<sup>-1</sup>: Ciomartan *et al.*, 1996; Madsen and Weaver, 1998) as well as mendipite, Pb<sub>3</sub>O<sub>2</sub>Cl<sub>2</sub> (at 330 cm<sup>-1</sup>: Bouchard and Smith, 2003). Thus, the bands at 318 and 332 cm<sup>-1</sup> in the Raman spectrum of napoliite should be assigned to stretching vibrations of the longer and the shorter Pb–O bonds, respectively (see Table 6).

Bands with the largest Raman shift of 202 cm<sup>-1</sup> are observed in the Raman spectra of lead chlorides cotunnite, PbCl<sub>2</sub>, and challacolloite, KPb<sub>2</sub>Cl<sub>5</sub>. However, these bands correspond to overtones (*e.g.* the strongest fundamental stretching band of cotunnite is observed at about 100 cm<sup>-1</sup>: Bouchard and Smith, 2003). Raman spectra of fluorocronite, PbF<sub>2</sub> (Krishnamurthy and Soots, 1970) and matlockite, PbFCl (Bouchard and Smith, 2003) contain distinct bands in the range 220–260 cm<sup>-1</sup>. These bands are absent in the Raman spectrum of napoliite. Consequently, bands below 200 cm<sup>-1</sup> in Figure 3 can be tentatively assigned to mixed soft lattice modes involving both Pb–F and Pb–Cl bonds as well as Cl–Pb–O and F–Pb–O angles.

Bands in the range 590–700 cm<sup>-1</sup> and at 447 cm<sup>-1</sup> in the IR spectrum of fiedlerite-1A, Pb<sub>3</sub>Cl<sub>4</sub>F(OH)·H<sub>2</sub>O are assigned to the Pb···O–H bending and H<sub>2</sub>O libration vibrations (Zubkova *et al.*, 2019). Thus, the absence of bands above 340 cm<sup>-1</sup> in the Raman spectrum of napoliite is in agreement with the conclusion on the absence of OH groups and H<sub>2</sub>O molecules in this mineral.

## **Chemical Composition and Chemical Properties**

Seven electron-microprobe analyses were carried out with a JEOL JXA 8230 Superprobe (WDS mode with an accelerating voltage of 20 kV, a beam current on the specimen of 20 nA and a beam diameter of 2 μm). Peak counting times (CT) were 20 s for all elements; CT for each background was one-half of the peak time. The raw intensities were converted into concentrations using *X-PHI* (Merlet, 1994) matrix-correction software. Both the crystal structure data and Raman spectroscopy confirm the absence of H<sub>2</sub>O and OH, borate and carbonate groups in the mineral. The contents of other elements with atomic numbers higher than that of carbon are below detection limits. Analytical data and used standards are given in Table 2.

The empirical formula calculated on the basis of 3 anions is  $Pb_{1.999}O_{0.997}F_{0.996}Cl_{1.007}$ . The ideal formula is  $Pb_2OFCl$ , which requires PbO 92.07, F 3.92, Cl 7.31,  $-O \equiv F+Cl$  -3.30, total 100 wt.%.

## X-ray diffraction data and crystal structure

The powder X-ray diffraction (XRD) data (Table 3) were collected with a Rigaku R-AXIS Rapid II diffractometer equipped with cylindrical image plate detector using the Debye-Scherrer geometry (d = 127.4 mm), Co $K\alpha$  radiation (rotating anode with VariMAX microfocus optics), 40 kV, 15 mA, and an exposure time of 15 min. The angular resolution of the detector is 0.045 20 (pixel size 0.1 mm). The data were integrated using the software package Osc2Tab (Britvin *et al.*, 2017). The unit-cell parameters refined from the powder data using UNITCELL software by Holland and Redfern (1997) are as follows: napoliite is tetragonal, space group  $P4_2/mcm$ , a = 5.737(1), c = 12.573(3) Å, V = 413.8(2) Å<sup>3</sup> and Z = 4.

To obtain the single-crystal XRD data a thin lamellar crystal of napoliite was mounted on a thin glass fiber and examined using Bruker APEX II DUO X-ray diffractometer with a Mo-I $\mu$ S microfocus X-ray tube ( $\lambda$  = 0.71073 Å) operated at 50 kV and 0.6 mA. More than a hemisphere of three-dimensional XRD data was collected with frame widths of 0.5° in  $\omega$ , and a 120 s count time for each frame. Then the collected data were integrated and corrected for absorption using a multi scan type model implemented in the Bruker programs APEX and SADABS (Bruker-AXS, 2014). The unit-cell parameters of napoliite [a = 5.7418(11), c = 12.524(4) Å, V = 412.9(2) Å<sup>3</sup> and Z = 4 in  $P4_2/mcm$ ] were determined and refined by least-squares techniques. The crystal structure was refined using SHELXL (Sheldrick, 2015) to  $R_1$  = 0.024 for 222 reflections with  $F > 4\sigma(F)$ . Crystallographic data and refinement parameters are reported in Table 4; coordinates and displacement parameters of atoms in Table 5 and selected interatomic distances in Table 6. The crystallographic information file has been deposited with the Principal Editor of *Mineralogical Magazine* and is available as Supplementary material (see below).

All attempts to refine the structure of napoliite in orthorhombic space groups, as reported earlier for synthetic [Pb<sub>2</sub>OF]Cl (Aurivillius, 1977), resulted in physically unrealistic displacement parameters for Pb, Cl and O atoms and instability of the refinement.

The structure of napoliite (Fig. 4a) contains one symmetrically unique Pb position. The Pb<sup>2+</sup> cation is coordinated by one O, one F, two mixed O/F sites and four Cl atoms (Table 6). The coordination geometry of the Pb atom in napoliite can be described as a distorted square antiprism. In agreement with previous results on Pb oxychlorides, the general feature of the Pb<sup>2+</sup> coordination in napoliite is the presence of several short Pb–X (X = O, F) bonds [2.2915(6)–2.5286(7) Å] located in one coordination hemisphere of the Pb<sup>2+</sup> cation. In the opposite hemisphere, the Pb<sup>2+</sup> cation forms four long Pb–Cl bonds [3.251(2)–3.566(2) Å]. This distortion is interpreted as the influence of the stereoactivity of  $s^2$  lone electron pair on the Pb<sup>2+</sup> cation.

Cl atoms have cubic coordination, whereas all *X* sites have tetrahedral coordination (Siidra et al., 2008), thus being central for anion-centered OPb<sub>4</sub>, FPb<sub>4</sub> and (O/F)Pb<sub>4</sub> tetrahedra (Fig. 4a). XPb<sub>4</sub> tetrahedra in napoliite are characterized by typical bond-length values (Table 6 and Fig. 4a). In napoliite, O1 atom forms four O-Pb bonds of 2.2915(6) Å each which is in a good agreement with the average bond length value of 2.33 Å in ideal OPb<sub>4</sub> tetrahedra observed for well refined structures (Krivovichev et al., 2013). The mixed oxygen/fluorine O/F1 site is tetrahedrally coordinated by four Pb-(O/F) bonds of 2.4129(5) Å, which is almost identical with the average Pb-(O/F) bond length of 2.41 Å in rumseyite (Turner et al., 2012). The presence of a site occupied by F in the crystal structure of napoliite is confirmed by the Pb-F bond length of 2.5286(7) Å. Anion-centered FPb<sub>4</sub> tetrahedra are also present in the structure of grandreefite, Pb<sub>2</sub>(SO<sub>4</sub>)F<sub>2</sub> (Kampf, 1991). The average F-Pb bond length in its structure is 2.55 Å.

## **Discussion**

Napoliite is a new representative (Fig. 4a) of a family of layered lead oxychlorides with lead oxide blocks derived from litharge PbO (Fig. 4b) (Boher *et al.*, 1985; Chukanov *et al.*, 2019). The structure of napoliite shows that the litharge-derived blocks may have different degrees and arrangements of F substitutions for O.

Napoliite is dimorphous with rumseyite (Turner *et al.*, 2012). Both minerals have not only identical chemical composition but also many similar physical properties. It should be noted that some properties of rumseyite (density, mean refractive index) were erroneously calculated by Turner *et al.* (2012); we use this occasion to correct these data (Table 7).

However, the crystal structures of these minerals, their space groups and powder X-ray diffraction patterns are different (Table 7, Fig. 4). Their origin is also different. Rumseyite is a secondary mineral formed due to the decomposition of galena in manganese oxide pods distributed in Carboniferous Limestone at the Torr Works (Merehead) Quarry in England, whereas napoliite is a fumarolic mineral deposited directly from hot gas as a volcanic sublimate. Both minerals can be considered as extremely rare; they have been found in a single specimen each and neither has a known, reliably characterised synthetic analogue.

From the crystal chemical point of view, a specific feature of rumseyite, [Pb<sub>2</sub>(OF)]Cl, is that all O<sup>2-</sup> anions in the [PbO] block are partially substituted by F<sup>-</sup> (Fig. 4 $\frac{1}{6}$ ). In synthetic orthorhombic Pb<sub>2</sub>OFCl (Aurivillius, 1977), F and O atoms are completely ordered over two symmetrically independent anionic sites (Fig. 4 $\frac{1}{6}$ ). Thus, napoliite represents a new structure type with a unique order/disorder pattern of fluorine and oxygen atoms. Its structural formula is [Pb1]Pb4]OO/F1(OF)F1F1Cl2, where litharge-derived block is in square brackets.

## Supplementary material

To view supplementary material for this article, please visit: https://doi.org/...

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**Table 1**. Reflectance data (R, %) of napoliite. The values required by the Commission on Ore Mineralogy are given in bold.

R <sub>max</sub> (%)	$R_{\min}$ (%)	λ (nm)	R <sub>max</sub> (%)	$R_{\min}$ (%)	λ (nm)
15.8	14.8	400	15.1	14.0	560
15.7	14.7	420	14.9	13.9	580
15.4	14.4	440	14.9	13.8	589
15.3	14.3	460	14.8	13.6	600
15.3	14.3	470	14.7	13.4	620

15.3	14.3	480	14.7	13.3	640	
15.3	14.3	500	14.7	13.3	650	
15.3	14.3	520	14.6	13.2	660	
15.2	14.1	540	14.6	13.2	680	
15.2	14.1	546	14.4	13.1	700	

Table 2. Chemical composition of napoliite (wt.%, mean of seven analyses).

Constituent	Wt.%	Range	S.D.	Standard
PbO	91.71	91.49–91.85	0.12	РЬТе
F	3.89	3.71–4.07	0.12	MgF <sub>2</sub>
C1	7.34	7.27–7.40	0.04	NaC1
-O = F + C1	-3.30			<b>5</b>
Total	99.64			

S.D. – standard deviation

**Table 3**. Powder X-ray diffraction data (d in Å) of napoliite. The eight strongest lines are given in bold.

$I_{ m obs}$	$d_{ m obs}$	$I_{\mathrm{calc}}*$	$d_{ m calc}**$	h k l
12	6.28	11	6.262	002
6	5.74	4	5.742	100
85	3.860	51	3.862	111
20	3.139	13	3.131	004
100	2.914	100	2.911	113
63	2.866	45	2.871	200
2	2.610	0.5	2.610	202
2	2.563	1	2.568	210
3	2.514	2	2.515	211
4	2.191	4	2.187	213
19	2.118	27	2.116	204
9	2.094	5	2.087	006
19	2.027	21	2.030	220
2	1.929	2	1.931	222
2	1.915	2	1.914	300
10	1.795	10	1.797	311
4	1.704	8	1.703	224
9	1.692	13	1.688	206
20	1.665	30	1.665	313
23	1.642	13	1.637	117
1	1.593	2	1.592	320
5	1.571	1	1.566	008

1	1.487	1	1.488	323
2	1 470	0.5	1.470	315
2	1.472	1	1.468	217
4	1.458	5	1.455	226
4	1.435	5	1.435	400
1	1.421	1	1.419	324
1	1.399	0.5	1.399	402
4	1.378	1	1.384	411
4	1.376	2	1.374	208
2	1.345	2	1.346	331
6	1.321	3	1.321	413
0	1.321	2	1.316	119
2	1.305	3	1.305	404
4	1.284	6	1.287	333
4		5	1.284	420
8	1.277	10	1.274	317

Table 4. Crystallographic data and refinement parameters for napoliite.

Crystal data	
Space group	$P4_2/mcm$
Unit cell dimensions $a, c$ (Å)	5.7418(11), 12.524(4)
Unit-cell volume (Å <sup>3</sup> )	412.9(2)
Absorption coefficient (mm <sup>-1</sup> )	81.909
Crystal size (mm)	$0.05 \times 0.05 \times 0.005$
Data collection	
Temperature (K)	<b>293</b>
Radiation, wavelength (Å)	Mo $K$ α, 0.71073
$\theta$ range (°)	3.548 - 27.971
h, k, l ranges	$-7 \rightarrow 6, -7 \rightarrow 7, -15 \rightarrow 16$
Total reflections collected	2961
Unique reflections $(R_{int})$	300 (0.05)
Unique reflections $F > 4\sigma(F)$	222
Structure refinement	
Refinement method	Full-matrix least-squares on $F^2$
Weighting coefficients a, b	0.0417, 0.196
Data/restraints/parameters	300/0/20
$R_1 [F > 4\sigma(F)], wR_2 [F > 4\sigma(F)]$	0.024, 0.063
$R_1$ all, $wR_2$ all	0.035, 0.070
Gof on $F^2$	1.090
Largest diff. peak and hole $(e \text{ Å}^{-3})$	2.479, -1.816

<sup>\*</sup>For the calculated pattern, only reflections with intensities ≥0.5 are given.

\*\*For the unit-cell parameters obtained by single-crystal X-ray diffraction.

Table 5. Coordinates and displacement parameters (Å) of atoms in napoliite.

Site	Wyck. site	х	у	Z	$U_{eq}$	$U^{11}$	$U^{22}$	$U^{33}$	$U^{23}$	$U^{13}$	$U^{12}$
Pb1	80	0.73266(7)	0.73266(7)	0.35353(4)	0.0180(3)	0.0190(3)	0.0190(3)	0.0160(4)	-0.00054(10)	-0.00054(10)	-0.0094(2)
C11	4i	0.7348(5)	0.2652(5)	1/2	0.0287(12)	0.0244(15)	0.0244(15)	0.037(4)	0	0	0.013(2)
F1	2b	0	0	1/4	0.019(7)	0.010(5)	0.010(5)	0.04(2)	0	0	0
O/F1*	4 <i>e</i>	0	$\frac{1}{2}$	1/4	0.018(6)	0.022(7)	0.026(7)	0.005(18)	0	0	0
O1	2 <i>d</i>	1/2	1/2	1/4	0.014(8)	0.008(6)	0.008(6)	0.03(2)	0	0	0

<sup>\*</sup>O<sub>0.5</sub>F<sub>0.5</sub>

 Table 6. Selected interatomic distances (Å) in napoliite.

Pb1-O1	2.2915(6)
Pb1-O/F1	$2.4129(5) \times 2$
Pb1-F1	2.5286(7)
Pb1-Cl1	$3.251(2) \times 2$
Pb1-Cl1	3.566(2) ×2
I UI-CII	3.300(2) ^2

 Table 7. Comparative data for napoliite and rumseyite.

Minoral	NI 1004 -	D
Mineral	Napoliite Napoliite	Rumseyite
Ideal formula	Pb <sub>2</sub> OFCl	Pb <sub>2</sub> OFC1
Structural formula	[Pb <sub>4</sub> O(OF)F]Cl <sub>2</sub>	$[Pb_2(OF)]Cl$
Crystal system	Tetragonal	Tetragonal
Space group	<i>P</i> 4 <sub>2</sub> / <i>mcm</i>	I4/mmm
a, Å	5.7418(11)	4.065(1)
c, Å	12.524(4)	12.631(7)
V, Å <sup>3</sup>	412.9(2)	208.7(1)
Z	4	2
Strongest lines	3.860 - 85	6.306 - 17
of the powder X-ray	3.139 - 20	3.848 - 41
diffraction pattern:	2.914 - 100	2.923 - 100
d, Å $-I$ , %	2.866 - 63	2.875 - 68
	2.118 - 19	2.110 - 12
	2.027 - 19	2.049 - 10
	1.665 - 20	1.719 - 9
	1.642 - 23	1.680 - 14
Colour	Colourless	Pale orange-brown
Streak	White	White
Transparency	Transparent	Translucent
Tenacity	Brittle	Brittle
Cleavage	Perfect on {001}	Perfect on {100}
Density (calc.), g cm <sup>-3</sup>	7.797	7.838*
Mean refractive index (calculated at	2.10	2.106*
589 nm)		
COM G ( COM)	1 4 2 /1 5 2	14.6
COM reflectance values (%)**	14.3/15.3	14.6
470 nm	14.1/15.2	13.6
546 nm	13.8/14.9	13.4
589 nm	13.3/14.7	13.2
650 nm		
Source	This paper	Turner et al., 2012

<sup>\*</sup>calculated by us from the chemical composition and unit cell parameters of rumseyite given by Turner *et al.* (2012)

\*\* $R_{\text{min}}/R_{\text{max}}$  for napoliite, R for rumseyite

## **List of Figures:**

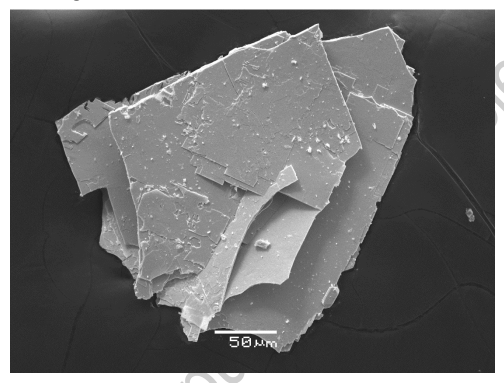


Fig. 1. Cluster of lamellar napoliite crystals. SEM (BSE) image. Sample # 5885/1.

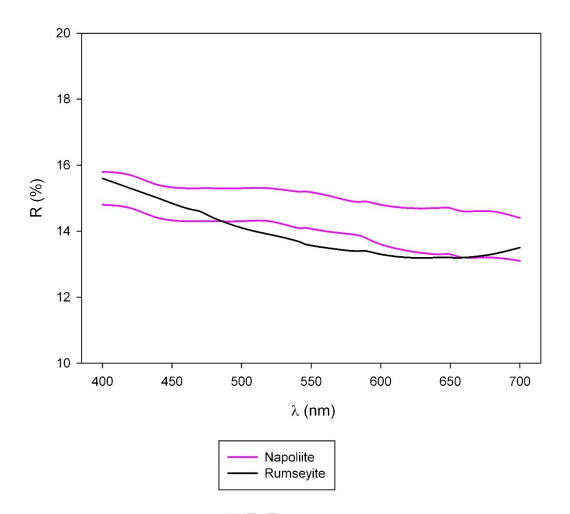


Fig. 2. Reflectivity curves for napoliite compared with the published data for rumseyite (Turner et al., 2012).

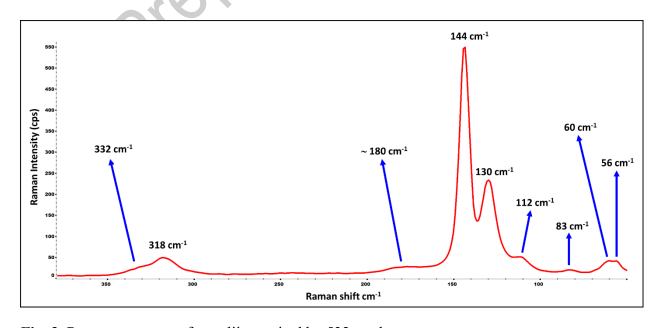


Fig. 3. Raman spectrum of napoliite excited by 532 nm laser.

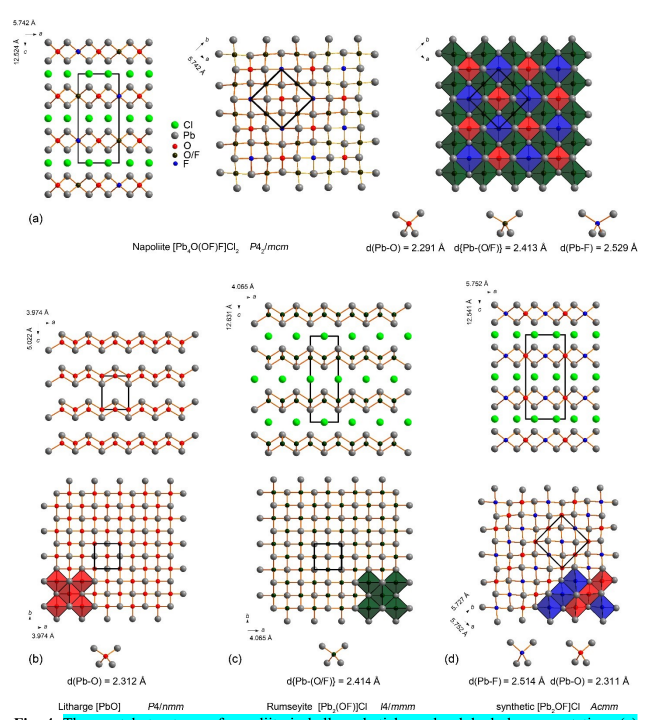


Fig. 4. The crystal structures of napoliite in balls-and-sticks and polyhedral representations (a) and structurally related litharge (Boher *et al.*, 1985) (b), rumseyite (Turner *et al.*, 2012) (c) and synthetic [Pb2OF]Cl (Aurivillius, 1977) (d). In structural formulae, litharge-derived blocks are given in square brackets. Average Pb-X (X = O, F) bond lengths in XPb<sub>4</sub> anion-centered tetrahedra are given for each crystal structure. Designations: grey balls = Pb, green balls = Cl, red balls = O sites, blue balls = F sites, dark green = mixed O/F sites; OPb<sub>4</sub> terahedra = red, (O/F)Pb<sub>4</sub> tetrahedra = dark green, FPb<sub>4</sub> = blue. Pb-Cl long bonds are omitted for clarity.