

Laphamite, an arsenic selenide analogue of orpiment, from burning anthracite deposits in Pennsylvania

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ABSTRACT. Laphamite, ideally $\text{As}_2(\text{Se,S})_3$, is monoclinic, $P2_1/n$, with $a = 11.86(1)$, $b = 9.756(9)$, $c = 4.265(9)$ Å, $\beta = 90.17^\circ$, $V = 493.5$ Å³, $Z = 4$, D (calc.) = 4.60, D (meas.) = 4.5(1) g/cm³. Microprobe analysis yielded: As 47.0, Se 43.7, S 8.7, sum = 99.4 wt. %. In plane polarized light in polished section at about 3200 K laphamite is moderately birefractant from white to grey. Reflection pleochroism is absent but fiery red internal reflections are characteristic as are golden yellow reflections along scratches. Anisotropy is moderate with grey rotation tints, masked in some sections by internal reflections. Reflectance spectra in air and oil between 400 and 700 nm are given. Colour values for the CIE illuminant C in air are, for R_1 followed by R_2 : $Y\%$ 29.0, 33.5, λ_d 480, 478; $P_e\%$ 9.1, 8.7. Laphamite forms dark red crystals, up to 5.0 mm, associated with orpiment, arsenolite, and an unnamed phase, NH_4AlF_6 , in a burning coal waste-dump at Burnside, Northumberland County, Pennsylvania.

KEYWORDS: laphamite, new mineral, orpiment, Burnside, Pennsylvania.

SECONDARY minerals formed as a result of burning coal seams or coal waste-dumps have been found in various parts of Pennsylvania, and in other places, for many years. The mineralogy of some of these occurrences was described in detail by Lapham *et al.* (1980) and a new mineral, downeyite, tetragonal SeO_2 , was described from such burning waste-dumps by Finkelman and Mrose (1977). Our investigation of a sample from dumps at Burnside, Pennsylvania, has resulted in

the description of another new mineral, $\text{As}_2(\text{Se,S})_3$. We have named this new mineral laphamite in honour of the late Dr Davis M. Lapham (1931-74), former Chief Mineralogist of the Pennsylvania Geological Survey (Socolow, 1976). This is a fitting phase to name for Dr Lapham inasmuch as he helped to initiate the study of the mineralogy of these coal dumps, and was particularly interested in the mineralogy of selenium (Barnes and Lapham, 1972; Lapham, 1975). The new mineral and the name have been approved by the Commission on New Minerals and Mineral Names, IMA. Type material is preserved at the Smithsonian Institution under catalogue No. 163039, and at the British Museum (Natural History) in a polished mount, E.1036, BM 1984,843.

Crystallography. Laphamite was studied using single-crystal Weissenberg and precession methods, and found to be monoclinic, $P2_1/n$, with unit cell parameters $a = 11.86(1)$, $b = 9.756(9)$, $c = 4.265(9)$ Å, $\beta = 90.17(30)^\circ$, $V = 493.5$ Å³, with $Z = 4$. This setting of the unit cell and space group is chosen for compatibility with those of orpiment, with which laphamite is inferred to be isostructural. The unit cell parameters were refined with powder data (Table I) obtained using a 114.6 mm diameter Gandolfi powder camera, a polycrystalline sample, Cu-K α X-radiation, and NBS Si as an internal standard. However, the β angle could not be refined because reflections for which d is a function of β

Table I. X-ray powder diffraction data for laphamite

I/I_0	d(Obs)	d(Calc)	hkl	I/I_0	d(Obs)
5	7.51	7.53	110	10	1.751
70	4.87	4.88	020	1	1.727
30	4.02	4.02	101	50	1.709
		4.01	101	5	1.686
2	3.90	3.91	011	2	1.598
40	3.72	3.71	111	1	1.571
		3.71	11 $\bar{1}$	1	1.559
10	3.22	3.21	021	1	1.528
20	3.10	3.10	12 $\bar{1}$	20	1.507
		3.10	12 $\bar{1}$	5	1.485
60	2.905	2.904	30 $\bar{1}$	2	1.429
		2.895	301	2	1.414
100	2.833	2.821	22 $\bar{1}$	1	1.399
		2.826	22 $\bar{1}$	2	1.332
80	2.773	2.783	31 $\bar{1}$	5	1.168
		2.775	311		
30	2.587	2.586	031	5	1.151
40	2.516	2.495	32 $\bar{1}$	5	1.139
		2.490	321	1	1.087
		2.512	330		1.074
5	2.439	2.439	040		
30	2.366	2.365	41 $\bar{1}$		
		2.359	411		
5	2.305	2.305	510		
1	2.255	2.256	240		
5	2.179	2.181	42 $\bar{1}$		
		2.176	421		
30	2.138	2.133	520		
		2.133	002		
20	2.081	2.085	141		
		2.039	141		
5	1.934	1.927	122		
		1.929	122		
		1.937	610		
5	1.919	1.916	530		
2	1.861	1.854	222		
		1.865	341		
		1.868	341		
50	1.777	1.774	051		
		1.783	032		

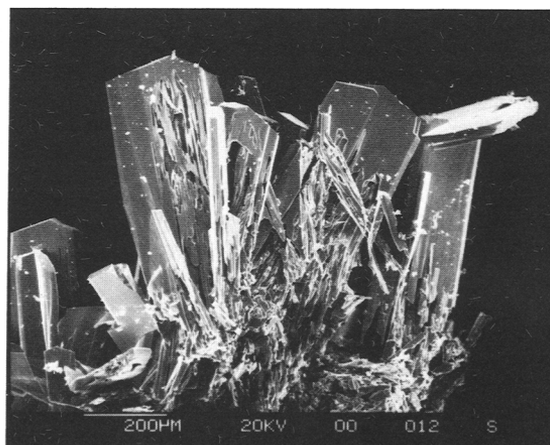
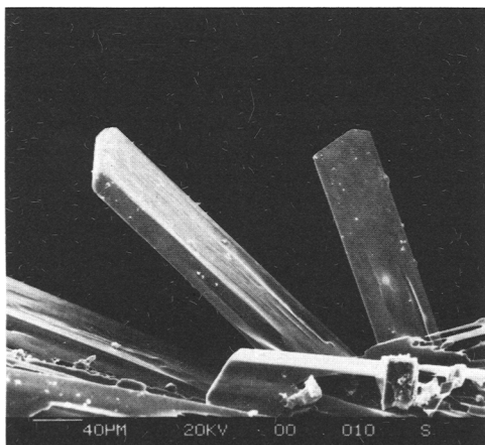
*Some d(Calc) values eliminated which have weak intensities as observed on single-crystal photographs.

could not be indexed unambiguously. It was therefore held constant at the value determined by single-crystal methods during the refinement of other parameters.

Morphologically, laphamite crystals are tabular on (010), elongate on [100], and are composed of the forms {100}, {110}, {101}, and {301}. Representative SEM images of laphamite crystals are presented as figs. 1 and 2. Many crystals are partially formed, or resorbed, as is common in sublimated phases.

Chemical composition. Laphamite was chemically analysed utilizing an ARL-SEMQ electron microprobe with operating conditions as follows: operating current 15 kV, sample current 0.025 μ A, measured on brass; standards: synthetic As_2Se_3 (As,Se), synthetic CdS (S). A wavelength-dispersive microprobe scan indicated the absence of any other elements with atomic number greater than 11, in measurable quantities. The data were corrected using a modified version of the MAGIC-4 program. The resultant analysis yielded: As 47.0, Se 43.7, S 8.7, sum = 99.4 wt. %. A formula, calculated on the basis of total atoms = 20, yields: $As_8(Se_{7.64}S_{3.72}As_{0.64})_{\Sigma 12.00}$, or $As_2(Se,S)_3$, with Z = 4. Laphamite crystals have quite variable Se:S ratios, but in all of them Se is dominant.

Physical properties. Laphamite occurs as dark red euhedral crystals, but the colour is easily discernible only in the smallest crystals; those larger than 1.0 mm are very dark red and nearly opaque in visible light. Maximum crystal length is 5.0 mm. Laphamite has a red-orange streak, resinous lustre, perfect (010) cleavage, and is extremely flexible, but not elastic. Laphamite is soft, but the hardness could not be determined quantitatively: loads of 5, 15, 25, and 100 g all produced indentations that were not measurable, owing to extreme malleability of the crystals. The density, measured with a Berman balance and temperature correction, is



FIGS. 1 and 2. Photomicrographs of laphamite. FIG. 1 (left). Free-standing prismatic crystals. FIG. 2 (right). A cluster of crystals illustrating the common habit.

4.5(1) g/cm³, compared with the calculated value of 4.60 g/cm³.

Optical properties in reflected light. Several crystals were mounted in cold-setting epoxy resin and polished with diamond abrasives (6, 3, 1, $\frac{1}{4}$ μ m) on Hypocel-Pellon laps with Engis machines. Laphamite is exceptionally difficult to polish to a scratch-free surface; only one of four crystals prepared was suitable for reflectance measurement. This was buffed, by hand, with a slurry of MgO in distilled water immediately before measurement.

In plane-polarized light (colour temperature about 3200 K) laphamite is moderately bireflectant from white to grey and lacks reflectance pleochroism. It has strong, fiery red, internal reflections. Golden yellow internal reflections are also apparent along scratches. Between crossed polars these internal reflections mask any anisotropy on (010) sections. On (001) the anisotropy is moderate with uncoloured (greys) rotation tints. Bireflectance measurements were made using the equipment and procedure described by Criddle *et al.* (1983) except that $\times 8$ objectives adjusted to effective numerical apertures of 0.1 were used with a SiC (Zeiss, 472) standard. The reflectance spectra (fig. 3) show that the dispersion of the reflectance differs for the two vibration directions measured, indicating a slight difference in colour. The dispersion is unchanged in oil (Zeiss N_D 1.515). Colour values calculated from the reflectance data (Table II) are similar for R_1 and R_2 in both media. The dominant wavelengths for the CIE A-illuminant (close to the colour temperature of the light sources used for the qualitative observations) suggest that the mineral should have a slightly bluish green hue. This would probably be

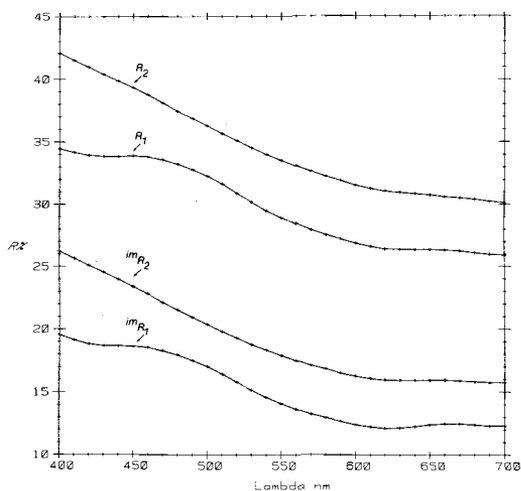


FIG. 3. Reflectance spectra in air and oil for laphamite.

TABLE II. Reflectance data and colour values for laphamite

λ nm	R_1	R_2	im_{R_1}	im_{R_2}
400	34.4	42.1	19.6	26.2
410	34.2	41.5	19.1	25.7
420	33.9	41.0	18.8	25.1
430	33.85	40.4	18.7	24.55
440	33.8	39.9	18.7	24.0
450	33.9	39.3	18.6	23.4
460	33.8	38.75	18.5	22.8
470	33.6	38.1	18.3	22.1
480	33.25	37.45	17.9	21.5
490	32.8	36.9	17.5	20.9
500	32.3	36.3	17.0	20.4
510	31.65	35.7	16.4	19.8
520	30.9	35.1	15.8	19.3
530	30.2	34.55	15.1	18.75
540	29.5	34.0	14.6	18.3
550	28.9	33.5	14.1	17.9
560	28.45	33.1	13.6	17.5
570	28.0	32.7	13.25	17.1
580	27.6	32.3	13.0	16.8
590	27.2	31.9	12.7	16.5
600	26.9	31.55	12.4	16.3
610	26.6	31.3	12.2	16.1
620	26.4	31.1	12.1	15.9
630	26.4	30.95	12.1	15.9
640	26.4	30.9	12.25	15.9
650	26.4	30.8	12.4	15.9
660	26.3	30.6	12.5	15.9
670	26.25	30.5	12.5	15.9
680	26.1	30.4	12.4	15.8
690	26.0	30.25	12.3	15.8
700	25.9	30.1	12.3	15.8

Colour values relative to CIE illuminant C				
x	0.290	0.292	0.277	0.281
y	0.300	0.298	0.286	0.285
z	29.0	33.5	14.15	17.9
λ_d	480	478	479	476
P_e	9.1	8.7	15.6	14.1

Colour values relative to CIE illuminant A				
x	0.428	0.430	0.415	0.420
y	0.403	0.402	0.397	0.396
z	28.3	32.85	13.6	17.4
λ_d	491	489	489	488
P_e	4.8	4.3	8.3	7.0

the case if the mineral were seen in association with others of similar luminance. As it is, the relative brightness of laphamite—8 to 10 times greater than the surrounding plastic—misleads the eye and the mineral is seen as uncoloured white-grey.

It was not possible to ensure that the vibration directions measured corresponded to principal sections of the crystal. For this reason, optical constants, calculated using the two-media method with the Koenigsberger equations, lack physical significance. Qualitatively, however, such calculations show that R_1 for laphamite is absorbing from 400 to 700 nm; furthermore, that R_2 becomes transparent between 490 and 620 nm. It follows that, in this region of the spectrum, as the reflectance is determined solely by the refractive index, the simpler Fresnel equations can be used with some confidence to derive the indices of refraction directly from the reflectances. Laphamite has an approximate n_{590} of 3.6.

Comparison of the R spectra with those published for orpiment by Caye and Padeloup (QDF,

1.6300, 1977), and Picot and Johan (1982, p. 283), reveal that R_1 for laphamite is similar to R_2 for orpiment, but R_2 is much higher—about 6% at 400 nm, and 10% at 700 nm.

Occurrence. Laphamite was discovered in a burning coal waste-dump at Burnside, Northumberland County, Pennsylvania, in the Western Middle Anthracite Field, approximately 1.5 km SSW of Shamokin, at 40° 46' 14" N., 76° 34' 12" W. It is one of the localities examined by Lapham *et al.* (1980) in their investigations. The specimen is a clinker, found as a surface encrustation around a single vent from which hot gases were being emitted (W. Downey, pers. comm.). Laphamite and massive, black, platy crystals (which give the X-ray powder pattern of NH_4AlF_4) are the dominant phases present. Octahedral crystals of arsenolite are found along the edge of some laphamite crystals. This association is illustrated in Fig. 35 of Lapham *et al.* (1980). Dark red amorphous gels, and a bright red soft granular aggregate which gives the powder pattern of orpiment, are also associated with laphamite. Crystals of laphamite were among the last to form, possibly by sublimation.

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