

THE CUMBERLAND FALLS, WHITLEY COUNTY, KENTUCKY, METEORITE.

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With analyses by EARL V. SHANNON, Assistant Curator.

This stone, which fell on the 9th of April, 1919, has been the subject of a note mainly descriptive of the fall,¹ by Prof. Arthur M. Miller, of Lexington, Kentucky, to whom the museum is indebted for his efforts in securing a considerable portion of the material.² The stone is of so unusual a type that it is worthy of more extensive notice than that given by Professor Miller, and fortunately the amount of the material secured is amply sufficient for the purpose.

On mere casual inspection there is little about the stone to suggest its ultra terrestrial nature. It is safe to say that had it not been seen to fall it would have been passed over by even those having a more or less intimate acquaintance with meteorites. On a broken surface it is of a light ash gray color, of a coarse texture, and might readily be mistaken for a terrestrial pegmatite in which the feldspar had undergone more or less whitening through weathering. Close examination reveals a pronounced brecciated structure (see pl. 15) produced by angular fragments of a chalky white mineral in pieces of all sizes up to 3 or more centimeters imbedded in a finer grey ground of apparently the same nature. Occasional inclosures of a dark gray-brown, almost black color, in one or two instances 3 to 5 cm. in diameter and angular in outline, exaggerate the pronounced brecciated structure which becomes so evident on a polished surface (pl. 15). Abundant flecks of a coal black, highly lustrous material are scattered irregularly through the ground, sometimes so abundant and of such small size as simply to render the rock dark gray in color, or again in shining blotches 10 to 15 mm. in diameter. Investigation shows these to be graphite. No iron or iron sulphide is noticeable on the broken surface and only abundant spots of newly formed iron rust suggest the presence of a ferrous chloride. The fusion crust or rind is inconspicuous, of a

¹ *Scienza*, June 6, 1919; also *The Mineral Resources of Kentucky*, vol. 1, ser. 5, No. 2, July 1919, pp. 110-114.

² Two complete individuals weighing respectively 567 and 2,347 grams (see pl. 14), and 13,476 grams of fragments. A 190-gram fragment was donated also by Mr. L. E. Bryant.

yellowish to dirty yellowish-brown color, smooth and extremely thin—a mere skin coating. In several instances there was noted on freshly broken surfaces, small, very thin areas of coal black or smoke-black glass. The cause of this or its relation to the crust is not readily apparent, but it is doubtless the very last fusion product of atmospheric resistance before reaching the earth. The usual pittings or thumb marks are present though the rock has become broken into so many pieces that these are not in all cases markedly evident. (Pl. 14.) A local slickensided movement is developed along the graphitic areas but which in no case observed extends throughout the mass. It would indicate merely such a movement as would take place within a mass under compression, without the production of faults. On the polished surface scattered particles of metallic iron and iron sulphide are readily observed, but they are extremely irregular in their distribution, and much more abundant in the dark, nearly black enclosures referred to. An interesting feature is the peculiar weathered appearance of even a fresh fracture. Fragments broken through the impact of fall and gathered within a few days show dead, lusterless surfaces, as though exposed for many weeks or months. It is probable that this is due to the physical condition of the main constituents, noted later.

In the thin section the white, chalky mineral referred to is seen to make up the main mass of the rock, though in various conditions of fragmentation from almost perfect forms to mere dust (pls. 16 and 17). These are often so crushed, crumpled, and otherwise distorted as to give only undulatory extinctions, and with other optical properties badly obscured. More perfect forms occur as broad plates (pl. 16, fig. 2) with well-defined vertical cleavage lines giving parallel extinctions. Basal sections show imperfect, nearly rectangular prismatic cleavage and the emergence of an optic axis. These facts together with the refractive indices (1.658+) and the results of Mr. Shannon's analyses (I, p. 100) leave no question but that the mineral is enstatite. In many sections, however, the mineral shows in polarized light irregular, wavy, and interrupted bandings which extinguish alternately as the stage is revolved, in a manner at first suggestive of the polysynthetic twinning of monoclinic pyroxenes or feldspars. In these cases the broader, more continuous bands give parallel extinctions and show in converged light the emergence of a bisectrix. The narrow, often indistinct and pinched out bands give inclined extinctions running as high as 37° . No distinction in color or refractive indices is noticeable, but there is apparently no question but there is here an intergrowth of orthorhombic and monoclinic forms ($\infty P \infty = \infty P \infty$) in the usual manner. If the analyses made by Mr. Shannon correctly represents this intergrown material (it was selected and analyzed before such an intergrowth was suspected) the proportional amount

of the monoclinic form as indicated by the amount of alumina and lime ($1.09 \text{ Al}_2\text{O}_3$ and 0.96 CaO) must be very small. The extreme narrowness of the bands giving the inclined extinction is, however, at least partially confirmatory of this.

The chalky appearance of the mineral is plainly due to its physical condition, an abnormal development of the cleavage, which incidentally causes it to crumble away under slight pressure and makes it susceptible of being ground to powder in an agate mortar as readily as so much calcite. Whether this condition is due to the shock which resolved the original mineral into fragments is uncertain, but it would seem most probable. The failure to become recompact under subsequent pressure might well be ascribed to a lack of moisture, pressure and dry heat alone naturally being less conducive to metamorphism.

In addition to the above, certain of the slides show intergrown with the enstatites in the form of small oval and irregular areas a brilliant polarizing mineral with the sharply developed platy structure characteristic of diallage. The mineral is also nearly colorless with a very faint green tinge, and gives extinction angles measured against the edges of the plates, that is, on clino-pinacoidal sections, as high as 27° . The proportional amount of the diallage is quite variable, some slides showing only an occasional rounded granule and others several of the intergrowths mentioned.

Scattered irregularly throughout the mass of the rock are the scale-like segregations of graphite above noted, sometimes several millimeters in diameter, in connection with which a differential movement has given rise to small areas with slickensided surfaces. In the finer portions, the graphite is so evenly and finely diffused as to impart a dark gray color. Metallic particles are quite inconspicuous excepting on a polished surface, as are also those of iron sulphide. The relatively greater abundance of the metal and sulphide in the dark inclosures above noted, is very evident on the polished surface (pl. 18). No calcium phosphate, maskelynite, oldhamite, osbornite, or other accessory minerals can be detected, although microchemical tests give rise to the usual globular ammonium-phospho-molybdate forms.

A close study of the dark inclusions developed some interesting and unexpected conditions. Examination with a pocket lens of the polished surface of one of the larger inclosures shown in plates 15 and 18 at once suggests a chondritic structure, a suggestion fully borne out by a study of the material in thin section, which shows a dark, obscure, and muddy ground containing numerous illy defined, compressed and distorted radiating, barred, and nearly holocrystalline chondrules of olivine and enstatite, mostly so obscured by a black impregnation that their true mineral nature is scarcely recognizable. In these respects the structure so closely resembles that of the McKinney and Travis

County, Texas, stones and others of Meunier's tadjerite group as to suggest a similar origin; that is, as developed from a normal chondrite (aumalite) through a process of heating. It is further to be noted that the dark portions are much richer in metal and, judging from the formation of abundant hydroxide of iron on a freshly cut or broken surface, richer also in ferrous chloride. These facts are borne out by the analyses noted later. The manner in which the metal occurs is interesting and peculiar, leaving no question as to its secondary origin and the foreign nature of the inclusion as well. One of these occurrences is shown enlarged some three diameters in plate 18, the metal in fine threads cutting across the surface in a manner strongly suggesting the figures sometimes given to show the play of lightning during a heavy electrical storm. Aside from these forms the metal at times completely surrounds a chondrule and even penetrates into it in the form of fine threads. The appearance is in entire accord with the idea of its late introduction after the crystallization.

It is evident at once that we have here a meteoric breccia composed of fragments of two quite dissimilar stones. This is sufficiently apparent from both megascopic and microscopic examination. The careful work of Mr. Shannon, the analyses quoted below, is fully confirmatory.

CHEMICAL ANALYSES BY E. V. SHANNON.

Before the intergrown nature of the pyroxenic constituents was suspected the clean, chalky-white portion was carefully sampled, crushed, and separated from possible impurities by the mercuric-iodide gravity solution. The results of an analysis of the powder thus obtained are given in column I below. In columns II and III are given for purpose of comparison previously reported analyses of enstatite from the meteorites of Bishopville, South Carolina, and Hvittis, Finland. The comparison with the enstatite of Hvittis, it will be noted, is particularly close.

Analyses of enstatite.

	Cumberland Falls.	Bishopville.	Hvittis.
	I	II	III
SiO ₂	59.53	59.97	59.05
Al ₂ O ₃	1.09		1.09
MgO.....	37.17	39.34	37.10
FeO.....	.98	.40	.90
CaO.....	.96		.98
(Ba, Sr)O.....	None.		Na ₂ O .68
NiO.....	None.		K ₂ O .47
Loss on ignition.....	.33		
Total.....	100.06	99.71	100.27

A 70-gram fragment which, so far as could be judged, was representative of the gray brecciated portion of the stone, was selected and, through the courtesy of Dr. George Steiger, ground at the laboratory of the U. S. Geological Survey. This yielded as in column I below. Unfortunately Mr. Shannon was not present during the process of grinding and it is possible that a larger portion of small particles of the dark stone were incorporated in the mass than was surmised from the appearance of the fragment. The probability of this, which was not at first realized, is suggested by the slight excess of magnesia (MgO) and ferrous oxide (FeO) in the bulk analysis over that in the enstatite given above. In columns II, III, and IV are given for comparison previously published analyses of the Busti, Bishopville, and Shalka stones. It will be noticed that so far as the magnesium is concerned the Cumberland Falls stone agrees very closely with that of the first-named, although a trifle higher in silica.

Bulk analyses of the light (major) portion of the meteorite.

	I	II	III	IV
Silica (SiO ₂).....	55.172	52.73	57.034	52.51
Alumina (Al ₂ O ₃).....	.382		1.706	.66
Chromic oxide (Cr ₂ O ₃).....	.062			1.25
Phosphoric oxide (P ₂ O ₅).....	Trace.			Trace.
Iron (Fe).....	.888		.181	.25
Manganese (Mn).....	.005			
Nickel (Ni).....	.059		.039	
Cobalt (Co).....	.004			
Copper (Cu).....	.003			
Chromium (Cr).....	Trace.			
Nickel oxide (NiO).....	.123	.78	.538	
Cobalt oxide (CoO).....	Trace.		Trace.	
Ferrous oxide (FeO).....	2.916	4.28	1.265	16.81
Lime (CaO).....	1.586	1.18	2.016	.89
Magnesia (MgO).....	38.734	37.22	33.506	28.35
Manganous oxide (MnO).....	.112	.01	.189	
Soda (Na ₂ O).....	.157		1.027	.22
Potash (K ₂ O).....	.150		.089	
Water (H ₂ O).....	.167	Ign.	1.995	
Sulphur (S).....	.784		.297	.14
Phosphorus (P).....	.034			
Chlorine (Cl).....	.028	¹ 2.35		
Carbon (C).....	.164	² .92		
	101.530	99.47	99.882	101.08
Less O for (Cl, S, P).....	.569		.147	
	100.961		99.735	

¹ Na₂S, CaSO₄, CaCl₂.

² By ignition.

The results given in column I seemingly bear out the microscopic determinations, and, in connection with the analysis of the white pyroxenic constituents given on page 100, warrant the conclusions

drawn as to the mineral composition of the stone. It is to be noted, however, that qualitative tests show an unusually large proportion of silicate matter soluble in acid, and suggest the need of further chemical work. This must, however, be deferred for the present.

The 0.888 per cent of metal yielded:

Iron (Fe).....	92.596
Nickel (Ni).....	6.152
Cobalt (Co).....	.417
Manganese (Mn).....	.522
Copper (Cu).....	.313
Chromium (Cr).....	Trace.
	100.000

Bulk analysis of the dark chondritic inclosure, yielded:

Silica (SiO ₂).....	41.683
Alumina (Al ₂ O ₃).....	1.537
Chromic oxide (Cr ₂ O ₃).....	.591
Ferrous oxide (FeO).....	9.399
Nickel oxide (NiO).....	.211
Cobalt oxide (CoO).....	trace
Phosphoric oxide (P ₂ O ₅).....	trace
Lime (CaO).....	4.059
Magnesia (MgO).....	27.848
Iron (Fe).....	12.108
Nickel (Ni).....	.747
Cobalt (Co).....	.078
Copper (Cu).....	.001
Chromium (Cr).....	trace
Manganese (Mn).....	.088
Potash (K ₂ O).....	trace
Soda (Na ₂ O).....	trace
Chlorine (Cl).....	.045
Sulphur (S).....	2.464
Phosphorus (P).....	.014
Carbon (C).....	.449
Ignition (H ₂ O).....	.210
	101.532
Less O for (Cl,S,P).....	1.448
	100.084

In comparison with the other chondritic stones this offers no unusual features.

Treated with dilute hydrochloric acid (sp. gr. 1.06) and sodium carbonate solution in the customary manner, the silicate portion, free from metal and metallic sulphide, yielded 22.582 per cent of soluble matter of the following composition:

Silica (SiO ₂).....	38.239
Alumina (Al ₂ O ₃).....	trace
Ferrous oxide (FeO).....	6.566
Nickel oxide (NiO).....	.043
Manganous oxide (MnO).....	.709
Cobalt oxide (CoO).....	trace
Lime (CaO).....	5.246
Magnesia (MgO).....	49.197
	100.000

The 56.58 per cent insoluble silicates yielded:

Silica (SiO_2).....	58.341
Alumina (Al_2O_3).....	2.705
Ferrous oxide (FeO).....	3.528
Nickel oxide (NiO).....	.295
Cobalt oxide (CoO).....	trace
Manganous oxide (MnO).....	.562
Lime (CaO).....	5.073
Magnesia (MgO).....	29.496
	100.000

This bears out the somewhat unsatisfactory determination of the prevailing orthorhombic nature of the pyroxenic constituent, but the high (5.073) per cent of lime (CaO) is difficult to account for.

The 13.022 per cent metallic portion yielded:

Iron (Fe).....	92.982
Nickel (Ni).....	5.735
Cobalt (Co).....	.599
Manganese (Mn).....	.676
Copper (Cu).....	.008
Chromium (Cr).....	trace
	100.000

The mineralogical composition of the dark inclusion as calculated from the foregoing is as below:

Metal.....	13.022
Troilite.....	6.760
Lawrencite.....	.080
Chromite.....	.869
Soluble silicates mainly olivine.....	22.582
Insoluble silicates mainly pyroxenes.....	56.580
Carbon, mainly amorphous.....	.449
Calcium phosphate.....	trace
Water, hygroscopic.....	.210
	100.552

The most striking features of the stone, aside from its coarse brecciated structure, are the marked evidences of compression manifested in the numerous small slickensided surfaces and the crushed and optically distorted condition of the pyroxenes, as shown both in the hand specimens and in thin sections. It is to be noted that while the original shattering which resulted in the production of the fragments may have been due to impact or explosive action, the mass has since been subjected to pressure under a heavy load whereby the particles have been further crushed and distorted and once more welded into a firm, rock-like mass. These are characteristics of deep seated terrestrial rocks that have been subjected to dynamic metamorphism.

The question naturally arises, is not the distortion so conspicuous in so much of the enstatite due to the crushing which resulted in the disintegration of the original meteorite rather than to any subsequent pressure? This question, I think, may be answered in the negative,

though not with absolute certainty. The study of the sections shows that the line of contact between the light stone and the dark inclusions, while apparently sharp, is, as shown in the thin section, quite irregular, as a rule particles from the one projecting into the other, though the superior hardness and toughness of the dark stone make this a less conspicuous feature than it might otherwise have been. Portions of the enstatite, are, however, jammed into the chondritic stone and particles of the chondritic stone into the enstatite as shown

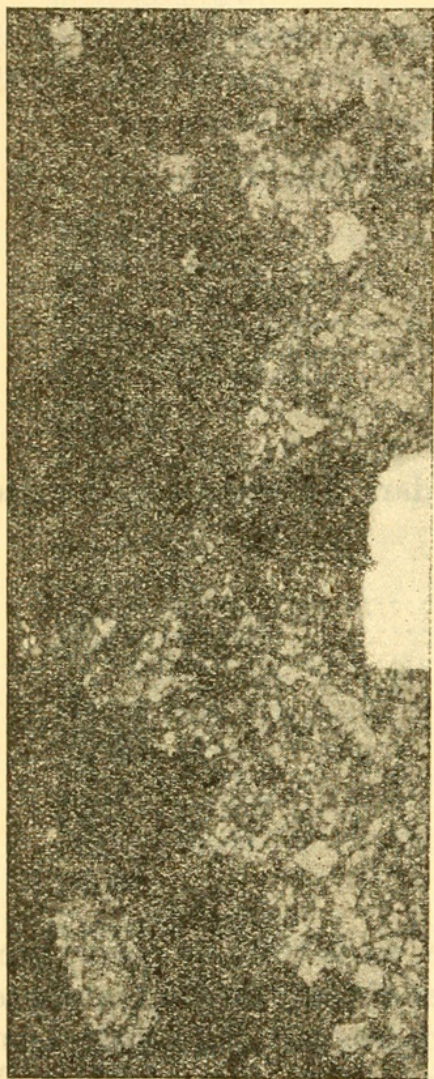


FIG. 1. SHOWING CONTACT BETWEEN DARK CHONDRITIC AND LIGHT ENSTATITE STONE.

in the accompanying figure. In one instance where a section has been so cut as to cause one of these interpenetrations to appear as an inclusion in the chondritic stone, a minute fault can be traced cutting through both pieces and making itself conspicuous by a slight off-set. Apparently, the admixture of the two kinds of fragments took place prior to the evident compression and both stones were involved. The numerous slickensided areas, sometimes of a few square centimeters dimensions, further testify to the compression and condensation in mass which the stone has undergone.

These structural characteristics are, as it seems to me, to be accounted for only on the supposition that the detrital matter composed of materials derived from the disintegration of previously consolidated rock masses of at least two distinct types, accumulated on the surface as in the case of an ordinary terrestrial volcanic breccia. Subsequently the beds were deeply buried and through crustal movements the material compressed into its present condition.

This carries with it the supposition that the meteorite is but a spawl from a very much larger mass, one of such size, indeed, as to have been subject to such crustal movements as are incidental to mountain making and which find their terrestrial counterpart in regions of maximum disturbance, as in the steep synclinal folds of our southern Appalachians. How large such a mass must be it is impossible to say, but that it must have been of planetary dimensions would seemingly be a safe assumption. In fact, that the fragments are direct evidence of the destruction of some preexisting planet seems a legitimate conclusion.

Incidentally, it may not be out of place to call attention to the fact that this adds one more to the most acidic type of magnesia-rich stones which have been seen to fall and all of which have come to us in a period of a little more than 100 years.¹

It must be evident from what has gone before that this stone has no exact counterpart among known meteorites and finds no exact place in the prevailing scheme of classification. Disregarding the inclosures of the chondritic stone it differs from the bustites, which chemically it closely resembles, in carrying no appreciable amount of oldhamite, plagioclase, or osbornite, and in its pronounced brecciated structure. From the chladnites it likewise differs in structure and its relatively high magnesia content. Nevertheless, it would seem more nearly related to these groups than others, though on the polished surface it suggests at first a remote similarity to the St. Michel stone described by Borgström and relegated by him to the Rhodites. The Cumberland Falls stone, however, carries no chondrules. It is a breccia, as already noted, and its mineral composition, aside from the chondritic inclusion, is limited almost wholly to the enstatite with an intergrown monoclinic form, sporadic diallage, and small quantities of metal, metallic sulphide, and graphite. In an attempt to make a position for it in the system of classification generally adopted,² I will suggest the name of *Whitleyite* (*Wht.*), and define it as a coarse white to gray breccia composed chiefly of enstatite with minor quantities of diallage, metal, metallic sulphide, and graphite, and with sporadic inclosures of a black chondritic stone. The term Cumberlandite might have been selected, but that this name has been preempted by Wadsworth³ for the terrestrial peridotite of Cumberland, Rhode Island. Whitley is the name of the county in which Cumberland Falls occurs.

EXPLANATION OF PLATES.

PLATE 14.

Two complete individuals showing crust and pittings. Actual sizes: 9.5 by 8.5 by 5 cm. and 17 by 16 by 7 cm.; weights, 567 and 234 grams, respectively.

PLATE 15.

Sawn and polished fragment showing brecciation and dark inclosures of chondritic stone. Actual diameters, 9 by 12 cm.

PLATES 16 AND 17.

Photo-micrographs under low magnification (about 5 diameters) showing structure.

PLATE 18.

Portion of slice shown in plate 15. Enlarged 3 diameters showing contacts between the fragmental enstatite and the chondritic stone and the peculiar distribution of the metal in the latter.

¹ See Merrill, G. P., The Percentage Number of Meteorite Falls and Finds considered with Reference to their Varying Basicity. Proc. Nat. Acad. Sci., vol. 5, pp. 37-39, February, 1919.

² See Wulffing, pp. 446-460.

³ Lithological Studies, p. 8.



Merrill, George P. 1920. "The Cumberland Falls, Whitley County, Kentucky, meteorite." *Proceedings of the United States National Museum* 57(2306), 97–105.
<https://doi.org/10.5479/si.00963801.57-2306.97>.

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DOI: <https://doi.org/10.5479/si.00963801.57-2306.97>

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