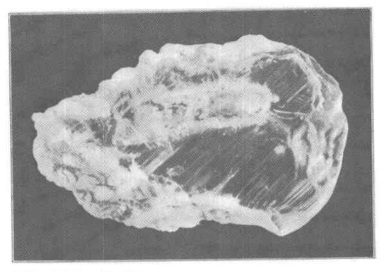


Front View



Photographs by Samuel & Gordon

Back View

GYPSUM CRYSTAL, ROUNDED BY WEATHERING, SHOWING A PHANTOM— A PSEUDOMORPH OF GYPSUM AFTER ANHYDRITE, STUDDED WITH QUARTZ. WEST PATERSON, N. J. NATURAL SIZE.

PLATE V

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A REVIEW OF THE GENESIS OF THE ZEOLITE DE-POSITS OF FIRST WATCHUNG MOUNTAIN, N, J.¹

VOL. I

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INTRODUCTION

NEW JERSEY is fortunate in having many famous mineral localities, and of these perhaps the most frequently visited by collectors are the trap quarries of First Watchung Mountain, especially those at West Paterson and Great Notch.

These occurrences have also received considerable attention from investigators, but the conclusions which have been arrived at as to the genesis of the minerals are not in general familiar to collectors. This is particularly true concerning some of the more recent discoveries, and the present paper has been written to call attention to these discoveries and the resulting conclusions as to the genesis of these remarkable mineral deposits.

Geology²

The rocks of this region are of Triassic age and consist of sedimentary rocks with interbedded basalt flows, the whole collectively known as the Newark Group.

The sedimentaries comprise fine-grained red shale, sandstone, conglomerate, and dark colored argillite or mudstone; the source of the materials having evidently been the gneiss, quartzite and limestone of the Highland belt to the west. Ripple-marks, mudcracks, rain-drop impressions, foot-prints of reptiles, and mineral cavities are frequently found. These sedimentaries are now believed to be of continental origin, deposited under semi-arid conditions in basins at the foot of the then newly uplifted Appalachian Mountains.

¹Read before the Philadelphia Mineralogical Society, September 14, 1916.

² See U. S. Geol. Survey, Passaic Folio, 157, 1911.

Numerous short but vigorous streams brought down the debris of the disintegrating and decomposing granites, gneisses and metamorphic sediments of earlier Paleozoic age, and deposited them in coalescing alluvial fans across the smoother plain of the crystalline Piedmont. Occasional downward movements of warping or faulting gave opportunity for local thickening of the deposits.

Similar deposits are now forming in the Great Basin of the western United States where extensive sediments are being laid down on the floors of sinking basins at the foot of the Sierras. In these saline residues, mud-cracks of great depth and ripplemarks are much in evidence. It is only under such conditions that thoro oxidation of the iron in sediments can occur; this, it is believed, accounts for the prevalent red color of the beds of the Newark Group.

In many of the beds glauberite⁴ and halite⁵ cavities are found, these minerals having formed in Triassic "playas" or temporary lakes, as they became concentrated thru evaporation.

During the deposition of these continental sediments, igneous activity interposed toward the close of the Triassic period. Three basalt flows mark this epoch in the vicinity of Paterson. Due to greater resistance to weathering as compared with the softer sedimentaries, the basalt masses rise above the surrounding country as prominent ridges, known as First, Second and Third Watchung Mountains. The mineral localities of importance are found in the first, those of West Paterson at a part of the mountain known as Garrett Rock.

The local features shown by the basalts are thus explained by Fenner.⁶

The flow spread over the surface of the accumulating sediments and filled the depressions occupied by the playa lakes. Over the dry areas, the lava came to rest quietly, and the normal process of cooling and crystallization followed without interruption. The rock formed under these conditions presents a dense, homogeneous mass, which, under the microscope, is found to have the holocrystalline texture normal to basalts. Over the lake beds, however, the lava . . . was quickly chilled from

³ J. Volney Lewis, The origin and relations of the Newark rocks, Ann. Rept. State Geologist of New Jersey, **1906**, p. 107.

⁴ E. T. Wherry, Glauberite crystal-cavities in Triassic rocks. Am. Min. (3), 37-43. ⁵ B. K. Emerson, Bull. U. S. Geol. Survey **126**, 144, 1895.

1, (3), 37-43. ⁶ B. K. Emerson, Bull. U. S. Geol. Survey 126, 144, 1895. ⁶ C. N. Fenner, Features indicative of physiographic conditions prevailing at the time of the trap extrusions in New Jersey, J. Geol. 16, 299, 1908. The Watchung basalt and the paragenesis of its zeolites and other secondary minerals, Ann. N. Y. Acad. Sci., 20, 93, 1910. the effects of water and vaporized steam. An exposure in the southwestern part of Paterson shows a mixture of lava and very fine mud several feet in thickness, and in many other places the lava was rendered extremely vesicular. The upper portions were not so thoroly impregnated with steam but were, nevertheless, quickly cooled and became viscous. The jets and tongues of fused material seem to have assumed the consistency of a thick syrup and instead of spreading laterally they solidified in smoothly rounded boulder-like masses, having considerable similarity to the 'pahoehoe' of Hawaiian flows. . . The rounded forms are sometimes built up to a thickness of 60–70 feet. The interior of the boulders cooled with sufficient slowness to permit the basalt to crystallize with normal texture, but each is sheathed with a crust of glass (tachylite) varying from an inch to several inches in thickness, having often a laminated structure. Where unaltered the color is usually dark olive-green or brown, and the appearance is decidedly vitreous. . .

The crusts frequently present a shattered appearance due to the sudden chill which they have experienced, and at times pockets among the boulders are filled with considerable masses of breccia of this nature.⁷

THE ZEOLITE DEPOSITS

Occurrence: The secondary minerals form nests and pockets in the angular spaces between adjacent boulder-forms and wrap around them in bands in the situation in which the glassy crusts were originally developed.⁸

This is due to the fact that the glass passed more easily into solution than the basalt, and formed the channels of easiest circulation.

. . Alteration also followed the shrinkage cracks which penetrate the interior of the normally crystallized interior of the boulders and produced veinlike bands of secondary minerals in a breccia of unaltered basalt.⁹

A study of the field relations has shown that the spheroidal masses of basalt, sheathed with glassy crusts, appear in areas which, at the time of the basalt flow, were covered by shallow lakes, and that these areas coincide with those in which the formation of zeolitic minerals has occurred.¹⁰

Source of the waters which deposited the zeolites: The query arises as to the origin of the circulating waters, whether meteoric or magmatic. . . In the case of these zeolitic deposits . . . several lines of evidence lead to the conclusion that the waters were of extraneous origin. Field observation shows that the secondary alteration extends to within a few feet of the lower surface of the trap sheet, and points to an uprise of waters from the underlying shales.¹¹

The aqueous contents of the lava would be completely expelled into the open air. The source of the waters which effected alteration must, therefore, be sought elsewhere. A theory of derivation from meteoric

⁷ Fenner, second paper cited, pp. 100 and 101. ⁸ Fenner, *ibid.* p. 109.

⁹ Fenner, *ibid.* p. 111. ¹⁰ Fenner, *ibid.* p. 102. ¹¹ Fenner, *ibid.* p. 104.

sources by a process of underground circulation appears to fill the requirements of the situation and to offer no obstacle. The preëxistent topographic depressions in the areas occupied by the lakes, altho buried by the flow of lava, would nevertheless be reached by waters flowing in from the edge of the sheet thru the porous sedimentary strata lying just beneath the impervious mass of normally crystallized igneous rock. Having reached such depressions, the hydrostatic head would be that due to the difference in elevation between the depressions and the point of ingress. The somewhat loose and porous structure of the accumulations of lava above the preëxistent lakes being favorable for percolation, the waters would naturally tend to rise to the surface. A circumstance favorable to this result lay in the fact that the lava retained a store of heat long after consolidation. Immediately after the latter event the temperature was so high that apparently all water which gained access was vaporized, notwithstanding the impediments to escape caused by restrictions of the channel; but as the temperature fell, the water level in this arm of the system of circulation slowly rose.

The system of circulation thus conceived is analogous to that of artesian flows, only modified in this respect, that the effective head was not so much due to a difference in elevation between the points of ingress and exit, as to vaporization of the water in the upper portions of the channel of exit.¹²

. . . Among the spheroidal masses, a considerable amount of interstitial space had been left. Moreover, the crusts were much shattered, and frequently the interiors of the masses were penetrated by a multitude of cracks produced by shrinkage in cooling. Where the openings were of sufficient size to permit free passage of superheated vapors from the water-impregnated sediments beneath, these gases seem to have rushed upward with great force and velocity, carrying with them quantities of finely comminuted dust from the lake beds and depositing it in the various interstices, in the form of a reddish-brown powder. . . . Mingled with this material, there are fragments of basaltic glass, whose nature is easily recognized.¹³

The general process was that of the slow but constant and uninterrupted cooling of igneous rock thru which aqueous solutions were percolating, deriving their heat from the enclosing rock and cooling as it cooled. . . With progressive fall of temperature conditions of equilibrium within the solutions were shifted, and new species were deposited. In these later changes, the material which participated was derived both from the first deposited minerals, which had now become unstable, and from new supplies of igneous rock, to which access was now gained.

The constant removal of material in solution undoubtedly enlarged the channels of circulations and, in places, produced cavities of considerable size. The most perfect crystals are naturally found in vugs of this kind. It does not appear, however, that such openings were at all necessary for recrystallization, and in most cases they were not present. The leaching appears to have been somewhat selective, for iron and magnesia are greatly reduced in the later stages.¹⁴

¹² Fenner, *ibid.* p. 106. ¹³ Fenner, *ibid.* p. 103. ¹⁴ Fenner, *ibid.* p. 112.

It was found difficult at first to account in a satisfactory manner for the introduction of such quantities of carbonic acid as the abundance of calcite requires. . . . The hypothesis which was finally adopted as being most probable was that the meteoric waters of the general form of circulation, . . . retained some store of the originally dissolved carbonic acid (probably as bicarbonate of lime, $CaH_2(CO_3)_2$, and carbonates of the alkalies) at the period of their history at which they entered the basalt sheet on the return journey to the surface. When conditions became favorable, this supply of dissolved carbonic acid reacted with the various original and secondary lime silicates at moderate temperatures. When the temperature is high, on the contrary, the process is reversed, so that carbonate rocks carrying silica, passing from the zone of katamorphism into the zone of anamorphism, where high temperatures prevail, have the carbonic acid driven off, while the lime combines with the silica, and produces lime silicates. The process is, therefore, easily a reversible one under variations of temperature. This probably accounts for the observed fact that little or no calcite appears to have been formed in the Watchung series during the prevalence of the more elevated temperatures under which the minerals of the first and second periods were deposited, and it was not until very moderate temperatures were reached that carbonation became the chief feature, tending to destroy the previously formed silicates.15

The theory of the source of the circulating waters above outlined is open to some question. It seems evident that the formation of the zeolites has locally been determined to a great extent by the development of basalt glass easily susceptible to alteration, which in turn was determined by the situation of the preëxisting lakes. Contact metamorphism of the shale beneath the basalt would render it quite impervious and an effective seal against invasion of waters from that direction. Even were such a circulation possible, it would undoubtedly have passed completely under the basalt to the opposite edge of the sheet, rather than thru the hot rocks above, as this would have formed the easiest path.

The present writer believes that the more plausible explanation is that the water involved was furnished both by the playa lakes and the magma itself. That considerable water was absorbed from these lakes by the basalt in its incursion cannot be doubted, and is evidenced by the development of anhydrite and glauberite, which distinguish these deposits from those of most other famous zeolite localities. Then, too, the solutions inherent in the magma cannot be disregarded.

At Moore Station, N. J., zeolites are found in contraction fissures in diabase, and are believed to owe their origin to mag-

¹⁶ Fenner, *ibid.* p. 175.

matic water exuded during the last stages of cooling into the contraction fissures. In the case of the Watchung basalt the zeolites may have been derived by alteration of the basalt glass and to a lesser extent of the normal basalt by solutions due to mingling of absorbed lake water and emanations from the cooling boulders.

The objection may be raised that the aqueous contents of the lava would have been completely expelled into the open air immediately upon its extrusion. Zeolites are, however, not uncommon in basalts, and amygdaloidal deposits are present in many occurrences of these rocks, and the origin in both cases may safely be ascribed to solutions from the original magma. It is quite possible that covering by later flows is a potent factor in the development of such zeolite deposits, for the cooling of the magma and formation of secondary minerals must under such circumstances continue long after other flows have sealed the top.

The mud found admixed with the zeolites and the glass would naturally be supposed to be the result of stirring up of the lake bottom by the basalt as it rolled along; much mud would be kept in suspension by circulating waters until the zeolites had been deposited,—often entering them as inclusions.

The majority of the secondary minerals are found to consist of the elements present in the original plagioclase feldspar, with or without addition of water. The possibilities of equilibrium among the great number of compounds present was considered in the light of Gibbs's phase rule, and some inquiry was made into the applicability of this rule to the conditions of the problem. It was deemed most probable that, altho various modifying conditions entered whose effect could not be wholly taken into account, the general conclusions to which the phase rule pointed should hold and that the possibility of all the compounds present coexisting in equilibrium could not be admitted.¹⁶

The processes of alteration may be divided . . . into . . . periods, which with their characteristic minerals are as follows:¹⁷

Period I. Boric acid period.

- Stage 1. Albite, quartz, garnet, amphiboles, specularite, sulfides.
- Stage 2. Datolite, prehnite, pectolite, amphiboles, specularite, sulfides.

Period II. Zeolite period.

Analcite, chabazite, heulandite, stilbite, natrolite, scolecite, laumontite, apophyllite, amphiboles, chlorite, specularite, sulfides.

Period III. Calcite period.

Thaumasite, calcite, gypsum, amphiboles, chlorite, specularite, sulfides.

¹⁶ Fenner, *ibid.* p. 186; see also pp. 115–121. ¹⁷ Fenner, *ibid.* p. 120.

Dr. Fenner has made an exhaustive study of the sequences of the minerals at West Paterson and Great Notch, examining many specimens and thin sections. The paper is replete with descriptions of slides and specimens, and extended discussions of the relationships of the minerals.

Attention may be called, however, to albite which is found replacing labradorite. Dr. Fenner believes that the various molecules entering into the feldspars are chemically independent, and presents the theory that

. . . the anorthite molecule was continually leached out of the labradorite, while albite was deposited in its place, a chief feature being probably a growth of crystals at the expense of others in the vicinity.^{18}

Some of the most interesting specimens to be found at the quarries are the well known "pseudomorphous cavities" found in quartz, prehnite, datolite, and pectolite. These have long been known under the misnomer "quartz pseudo pectolite" or in a few instances, quartz pseudo some zeolite. Two types of crystals are represented by the cavities, one rectangular in outline, thick to thin tabular in habit, and evidently orthorhombic; the other lozenge or "diamond"-shaped in cross section, prismatic in habit, and monoclinic in symmetry.¹⁹

Babingtonite was described from Paterson by Dr. Fenner²⁰ in 1914 and he suggested that this was the original mineral of the cavities. But Mr. F. I. Allen²¹ discovered anhydrite in McKiernan and Bergin's quarry about the same time, in all stages of alteration to gypsum or thaumasite; and study has shown that this mineral was undoubtedly the one to which most of the rectangular cavities are due.

More recently loose casts of quartz in the lozenge-shaped cavities have been found, showing well defined prismatic and pyramidal faces, and these were shown by Dr. Wherry²² to be due to glauberite.

¹⁸ Fenner, *ibid*. p. 122.

¹⁹ E. T. Wherry, The lozenge-shaped cavities in the First Watchung Mountain zeolite deposits, J. Wash. Acad. Sci., 6, (7), 182, 1916.

²⁰ C. N. Fenner. Babingtonite from Passaic County, N. J., J. Wash. Acad. Sci., 4, 553-558, 1914. Additional notes on babingtonite from Passaic Co., N. J., *ibid.*, 598-605.

²¹ F. I. Allen, N. Y. Times, Nov. 23, 1914; Am. J. Sci., 39, 134, 1915.
²² E. T. Wherry, loc. cit.

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An interesting specimen of gypsum is shown in Plate V. The back o the crystal shows a pseudomorph of gypsum after anhydrite. The anhydrite was evidently the first mineral to form, and received a coating of quartz crystals; later, the anhydrite was replaced by gypsum, which recrystallized and became the nucleus of a much larger crystal, which has since suffered corrosion of the faces as shown in the first figure.

Mineral	Composition					Genetic rela	lion	
Labradorite	1 Na Al			Si ₃ O ₈		Primary		
Labradorite	1-3 Ca Al			Si2Oa		44		
Babingtonite	(Ca, Fe, Mn) Fe ₂			$(SiO_3)_4$		6 6		
Augite	∫m Ca Mg			$(SiO_3)_2$		4.6		
Augite	\n (Mg. Fe) (Al, Fe)2			SiC ₆				
Olivine	(Mg, Fe)2			SiO4		4.4		
Albite	Na Al			Si3O8			al metamorph	
Amphibole	∫ (Ca, Mg, Fe)			SiOa		4.4	**	
Amphibole	1	Na Al		(SiO3)2			**	
Garnet	(Mg, Fe) ₃ Al ₂			(SiO4)3		6.6		
Chlorite	Ha (Mg, Fe) ₄ Al ₂			$(SiO_6)_3$		8.4	44	
Specularite		Fe2		O3		4.6	6.6	
Pyrite?		Fe		S_2		4 s	"	
Chalcopyrite		Fe Cu		S_2		11	**	
Sphalerite		Zn		s		4.4	••	
Galena		Pb		S		4.6	4	
Quartz				SiO ₂		+ 4	6 6	
Chalcedony				SiO_2		4.6		
Calcite		Ca		CO ₃		14	6.6	
Anhydrite		Ca		SO4		4.6	4.6	
(Glauberite)		Ca Na	1	(SO4)2		4.8		
Analcite		Na	Al	Si2Os.	H_2O	4.6		
Chabazite		(Na2, C	a) Al ₂	(SiOa)4.	$6H_2O$	66	12	
Gmelinite		(Na ₂ , C	a) Al_2	(SiOs)4.	6H ₂ O	4.6	- 440	
Pectolite	н	Na C		(SiOa)a		6.8		
Datolite		C	a B (OH)	SiO4		6.4	44	
Apophyllite	(H, K ₂) Ca			(SiO3)2.	H_2O	4.6	6.6	
Prehnite	H ₂	C	a ₂ Al	(SiO4)3		6.6	4.6	
Natrolite		Na ₂	Al2	Si3O10.	$2H_{2}O$	4.6	6.8	
Scolecite		Ca	Alz	Si3O10.	$2H_2O$		4.6	
	(m	Na ₂	Al2	Si3O10.	$2H_2O$	1.6	64	
Mesolite	ln	Ca	Alz	Si3O10.	2H ₂ O	**	4.6	
Thomsonite		Ca	Al_2	Si2O8.	$2\frac{1}{2}H_2O$	4.6	6.4	
Laumontite		Ca	Al2	(SiO3)4.	$4H_{2}O$	* 6	6.4	
Stilbite		Ca	Al_2	Si6O16.	$6H_2O$	* 6	4.4	
Heulandite		Ca	Al_2	Si6O16.	$5H_2O$	44	4.6	
Opal				SiO2.	nH2O	Weatherin	g product	
Gypsum		Ca		SO4.	$2H_{2}O$	44	**	
Thaumasite		Caa	(OH) ₃ (CO ₂)	(SO3) (H	(SiO4) - 12	2H ₂ O ''	46	
Aragonite		Ca		CO3		4.4	6.6	
Malachite		Cu2	(OH)2	CO ₃		* 5		
Azurite		Cua	(OH)2	(CO ₃) ₂		**	**	
Limonite		Fee	O3(OH)6			44	11	
Stevensite	H_2	Mgs		(SiO3)4.	H_2O	6.8	4.6	
Burgerite and r			ion products of		te of inde	finite composit	ion.	

TABLE I. PRINCIPAL MINERALS OF THE DEPOSITS