

90035A.1

Fayalite

Babson Farm Quarry

Rockport, Essex Co.,

Massachusetts

A.C. Gosse and G.M. Flint, Donors

MINERALOGICAL MUSEUM, HARVARD UNIVERSITY
CAMBRIDGE, MASSACHUSETTS

SPLITS

1. Max Cormen U. Huston
2. Alex Speer VPI & SU

HARVARD UNIVERSITY
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February 28, 1979

Dr. J. Alexander Speer
Department of Geological Sciences
Virginia Polytechnic Institute and
State University
Blacksburg, Virginia 24061

Dear Alex:

Egg on my face.

Well, now Uncle Paul has another amusing anecdote to tell in probe class. My spurious data resulted from probing a specimen that had not been carbon coated. In my excitement about having the Mg end-member, I transferred the grain mount for a sample holder which hadn't been coated to one which had. Hence all the samples except the forsterite gave reasonable results.

Nothing ever happened to the Austinville aragonite. It remains in the fluorescent exhibit in the museum labelled hydrozincite. A similar specimen was in the fluorescent exhibit in Dr. Michael's home.

A specimen of Rockport fayalite will follow. The synthetic tephroite I sent Paul should be an ideal standard for manganese. What other elements do you need?

Charlie's Picker is up. I'm supposed to get on it later in the week. A second data set will be collected on another Mg-Mn olivine later in the Spring.

Would you look into the question of what has become of the stack of computer output I left with JBH? It would be helpful for the next refinements. If you can locate it, please squirrel it away until I can come down later in the Spring.

I'm glad to hear that you are writing. Soon you'll be rich and famous! Will you and the Higgins be visiting here this summer?

Best regards,

Carl

Carl A. Francis
Curator

CAF:cep

* Please send me copies of analyses and other data generated for this specimen.

Etudes Sur la Partie Occidentale du Lac de Geneve Courants et Temperatures par A. Betant et G. Perrenoud. Atlas, 12 Planches. Vol. 41. Fascicule 2. Pp. 226-293; 34 figs. Memoires de La Societe de Physique et D'Histoire Naturelle De Geneve, 1932. Prix du Texte et de l'Atlas: Fr. 20.

Zoological Parks, Aquariums and Botanical Gardens; by L. C. Everard. Pp. 72. Publications of the American Association of Museum. Washington, D. C., 1932.

The Felidae of Rancho La Brea: John C. Merriam, Chester Stock. Carnegie Institution of Washington. Pp. xvi, 231; 42 pls., 152 figs.

The Action of the Living Cell; Fenton B. Turk. New York, 1933 (The Macmillan Co., \$3.50).

Dissection of the Cat; by Jacob Reighard and H. S. Jennings. New York, 1932 (Henry Holt and Co., \$1.25).

Carnegie Institution of Washington. Year Book. No. 31. 1932. Pp. xix, 392.

OBITUARY.

DR. ORMOND STONE, professor of astronomy at the University of Virginia and director of the Leander McCormick Observatory from 1882 to 1912, died on January 17 at the age of eighty-six years.

DR. ARTHUR GRAY LEONARD, professor of geology at the University of North Dakota, died on December 17 at the age of sixty-seven years.

DR. WILLIAM LISTENARD ROBB, since 1902 professor of electrical engineering at Rensselaer Polytechnic Institute, died on January 26 at the age of seventy-one years.

DR. HARLAN WILBUR FISK, of the department of terrestrial magnetism, Carnegie Institution, died on December 26 at the age of sixty-three years.

DR. WILLIAM ARNON HENRY, emeritus professor of agriculture at the University of Wisconsin, died on November 24 at the age of eighty-two years.

DR. VICTOR STERKI, assistant curator of zoology in the Carnegie Museum, Pittsburgh, died on January 25 at the age of eighty-six years.

SIR JOHN ARTHUR THOMSON, professor emeritus of natural history at the University of Aberdeen, died on February 12 at the age of seventy-two years. In addition to his zoological investigations, he published a number of valuable books on the popular side of biology.

DR. JAMES JOHNSTONE, the British zoologist and oceanographer, died on December 27 at the age of sixty-two years.

DR. MALCOLM EVAN MACGREGOR, of the entomological laboratories at Esher, Surrey, England, died recently at the age of forty-three years. He was early a Carnegie fellow at Harvard.

DR. CLAUDE METFORD THOMPSON, professor emeritus of chemistry at the University College, Cardiff, died on January 4 at the age of seventy-seven years.

AMERICAN JOURNAL OF SCIENCE

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THE SYSTEM, $\text{Ca}_2\text{SiO}_4\text{-Fe}_2\text{SiO}_4$.

N. L. BOWEN, J. F. SCHAIER, AND E. POSNJAK.

INTRODUCTION.

The compound, Fe_2SiO_4 , is found in nature as the mineral fayalite, known from a number of localities but not of common occurrence. The compound, Ca_2SiO_4 , has been found in only one locality as a natural mineral, viz., near Larne, County Antrim, Ireland, and has been named larinite.¹ Both minerals are formed only under rather special conditions and are not of great importance as rock-forming minerals, but a knowledge of their stability relations may well throw light on factors involved in the formation of more important minerals with which they are associated.

The system, $\text{Ca}_2\text{SiO}_4\text{-Fe}_2\text{SiO}_4$, represents a join, specifically the orthosilicate join, in the more general system, CaO-FeO-SiO_2 , upon the investigation of which we are now engaged. The results of a study of one of the fundamental systems of this general system, viz., the system FeO-SiO_2 , have recently been published.²

To metallurgists the join now under discussion is of considerable importance because slags having such compositions are frequently used and are termed by them, monosilicate or singulo-silicate slags, following a system of naming silicates that has fallen into disuse in mineralogy.

METHOD OF INVESTIGATION.

The method which we have found satisfactory for the investigation of silicates containing ferrous oxide is described in full detail in the paper referred to above. In brief it may be said that mixtures were made up by taking silica, ferrous oxalate (or Fe_2O_3) and calcium carbonate in the proportions calculated to give the desired product and melting them

¹ Tilley, C. E., *Min. Mag.*, 22, 77-86, 1929.

² Bowen and Schairer, this Journal, 24, 177-213, 1932.

together three times in a platinum crucible in a gas furnace. Thus was obtained a product deviating somewhat from the desired composition by reason of a slight deficiency in iron (taken up by the platinum crucible) and the existence of some of the iron in the ferric state. Small samples of the mixture were then heated at measured temperatures in crucibles of pure electrolytic iron in a stream of nitrogen and the temperatures of beginning of melting, completion of melting, and of any other changes of phase were determined by the method of quenching, combined with microscopic determination of the phases in the quenched product. Upon heating in the iron crucible, a small change in the composition of the charge occurs which is principally due to reduction of ferric oxide by the iron crucible and consequent increase of the total iron content of the charge. This increase may partly balance or more than balance the deficiency produced by melting in platinum but the extent of balance cannot be predicted. A sample of each mixture was, therefore, subjected to chemical analysis after heating in an iron crucible at the temperature at which a change of phase was found to occur. Ordinarily, the temperature of heating was that at which completion of melting occurs and the analysis thus gave the composition of the liquid at the liquidus point for that composition. By repeating this procedure for a series of mixtures and plotting the determined liquidus and solidus temperatures and analyzed compositions (with microscopic determination of the crystalline phases) the equilibrium diagram was constructed.

Chemical Analysis of Charges.

Ferrous iron was determined by the Pratt³ method modified somewhat. The exact procedure was as follows: Weigh 0.2 to 0.3 gram sample into a 150 cc. transparent silica glass⁴ Erlenmeyer flask, add about 60 cc. distilled water. Displace the air in the flask by a stream of CO₂ (free from air) entering by a silica glass tube through a two-hole rubber stopper. Heat to boiling, remove flame and add 15 cc. (1:1 by volume) H₂SO₄ and 5 cc. of 40% hydrofluoric acid from a platinum

³ This Journal, 48, 149, 1894.

⁴ Silica glass was used to secure the advantage of observing the progress of solution of the sample. Other glasses would meet this requirement but were found to affect the FeO determination seriously. Blanks run on the silica glass used showed that no material which affected the FeO determination was dissolved from the glass by the acids.

dish by just lifting the stopper while CO₂ is still running. Replace stopper and boil gently until sample is completely dissolved (3 to 8 minutes). Cool in ice bath for 15 minutes, with CO₂ stream still running. Stop CO₂, remove from ice bath and add a large excess of solid boric acid (about 5 or 6 grams), titrate at once with standard KMnO₄ solution.

Total iron was determined by dissolving 0.2 to 0.3 gram sample in 15 cc. (1:1 by volume) H₂SO₄ and 8 cc. of 40% hydrofluoric acid in a large platinum dish and heating on a hot plate until all H₂F₂ had disappeared. The dish was cooled, contents diluted, and 5 cc. of concentrated HNO₃ added and the liquid transferred to a 250 cc. beaker. The iron was separated from CaO by triple precipitation using a large excess of NH₄OH and washing the precipitates well with hot water. The precipitate was ignited and weighed as Fe₂O₃. The difference between total iron and FeO gave Fe₂O₃. To check the separation of iron from CaO by the method of triple precipitation, on one sample the total iron was determined by reduction with H₂S and titration with KMnO₄. The results were practically identical.

EXPERIMENTAL RESULTS.

The results of the determination by these methods of the temperatures at which changes of phase occur, together with the analyzed composition of the charges, are given in Table I.

TABLE I.

Melting temperatures of mixtures.

at liquidus temperature				Temp. of completion of melting	Solid phases in equilibrium with liquid at liquidus temperature
CaO	FeO	Fe ₂ O ₃	SiO ₂		
7.48	59.99	2.25	29.39	1205 ± 2°	Fayalite and Fe
12.95	53.80	2.43	30.17	1160 ± 2°	Ca-Fe olivine and Fe
14.97	51.57	2.61	30.85	1117 ± 3°	Ca-Fe olivine and Fe
22.47	43.53	2.54	31.46	1122 ± 3°	Ca-Fe olivine and Fe
29.75	35.80	2.55	31.90	1182 ± 2°	Ca-Fe olivine and Fe
33.80	31.17	2.80	32.25	1208 ± 2°	CaFeSiO ₄ and Fe
36.21	28.83	2.71	32.75	1226 ± 2°	Ca-Fe olivine and Fe
39.34	26.35	1.84	32.47	1300 ± 5°	Ca ₂ SiO ₄ and Fe
43.02	21.82	2.15	33.01	1425 ± 5°	Ca ₂ SiO ₄ and Fe
				1525° (appr'x.)	Ca ₂ SiO ₄ and Fe

The table shows that even after holding the mixtures in a molten condition in an iron crucible in nitrogen some of the iron of the liquid is in the ferric state. In our former publica-

tion it was shown that even when the mixtures are melted in iron in an evacuated, sealed, silica-glass tube there is still failure of complete reduction of the iron of the mixtures to the ferrous state and reasons were there given for the belief that the small amount of ferric iron persisting in our runs made in an iron crucible in nitrogen represents a true equilibrium relation with metallic iron. Iron is incapable of further reducing the ferric iron content of the liquid; indeed, an attempt at further reduction by a powerful reducing agent would result only in the separation of metallic iron from the liquid without decrease in its Fe_2O_3 content. We have not regarded it as necessary in the case of the present mixtures to resort to any of the special procedures that were adopted in the earlier work to prove this fact since it had already been proved for one mixture that is common to both systems, viz., a mixture of the composition of fayalite.

The manner in which the content of Fe_2O_3 varies with the total iron content of liquids of the present series in equilibrium with metallic iron is of considerable interest. In the work on the system, FeO-SiO_2 , it was found that the Fe_2O_3 content always decreased with content of total iron, though not at a uniform rate, the decrease being more rapid at high than at low iron concentrations. When the iron content is decreased by adding, not silica alone as formerly, but both silica and lime as in the present series, no simple decrease of Fe_2O_3 content is obtained. Indeed, as one passes from fayalite towards Ca_2SiO_4 , the amount of Fe_2O_3 in the liquid in equilibrium with metallic iron does not at first decrease but actually increases slightly. The amount of increase is within the limit of error of analysis of an individual mixture, but the trend of the series, representing a decrease of total iron content (expressed as FeO) from about 70 per cent to less than 35 per cent, is distinctly upward, being 2.25 per cent Fe_2O_3 at fayalite and about 2.8 per cent at a point more than half-way over to Ca_2SiO_4 . The maximum appears to come at the point where the solid phase in equilibrium with liquid (in addition to solid Fe) changes from Ca-Fe olivines to Ca_2SiO_4 . Beyond that point the Fe_2O_3 content of the liquid decreases, but we are unable to follow the decrease far on account of the fact that the mixtures melt at too high a temperature to obtain complete melting in an iron crucible. The Fe_2O_3 content must, of course, become zero when the total iron content becomes zero at Ca_2SiO_4 .

Ternary Diagram of Equilibrium.

On account of the fact that the Fe_2O_3 content of the liquids cannot be reduced to zero, a ternary diagram is necessary for the accurate presentation of the results. Such a diagram is given in Fig. 1. It shows that all the mixtures on the orthosilicate join lie in the field of metallic iron and that all the orthosilicate mixtures, if perfectly pure, would melt incongruently with separation of metallic iron. Not only is this true but even if some Fe_2O_3 is present in the crystalline orthosili-

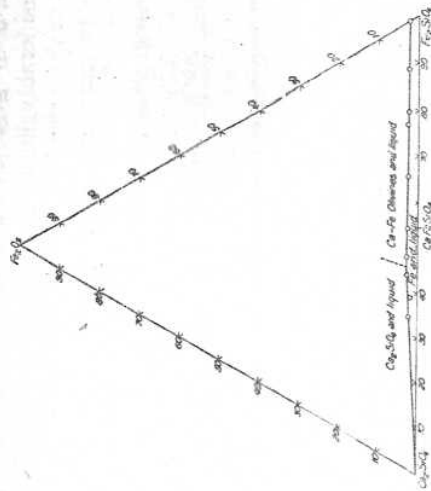


Fig. 1. Ternary diagram showing the composition of the liquids in equilibrium with metallic iron and crystalline orthosilicates of Ca and Fe.

cates its amount must exceed a certain minimum value in order to carry the composition outside the field of metallic Fe and eliminate the formation of metallic Fe upon melting. The minimum value for each orthosilicate composition is, of course, that given in Table I and shown by the position of the boundary curves limiting the field of iron in Fig. 1.

Incongruent Melting of Fayalite.

In our former paper we demonstrated that fayalite melts incongruently with separation of iron. For this purpose we used natural fayalite from Rockport, Massachusetts, and from Mourne Mountains, Ireland. Since then we have determined the chemical composition of the Rockport fayalite, the precise material that was used for our thermal investigation.

iron powder, Fe_2O_3 , and SiO_2 were mixed in the proper proportions, sealed in an evacuated silica-glass tube, and heated at 1100° for 4 days. In this manner a satisfactory product consisting of small, equant grains of fayalite, free from uncombined iron, was obtained. This material, after the microscopic examination which determined its character as just described, was again sealed in an evacuated silica tube and suspended in a furnace whose temperature was already fixed at 1255° , some 50° above the melting point of fayalite. Unfortunately it was found that the fayalite could not be held long under these conditions, for the liquid rapidly attacks the silica tube. In only $2\frac{1}{2}$ minutes some liquid had penetrated to the outside of the tube, so that runs of even moderate length, such as one might expect to be necessary to obtain the full separation of the equilibrium quantity of iron, were not practicable. Nevertheless, in one run in which the charge was left in the furnace only one minute and in which it had happened to be placed unsymmetrically in the furnace, it was found that the one side of the charge had melted and the other had not. The unmelted part remained in the same equant grains as the original material, the melted part recrystallized as bladed crystals. In the bladed portion rare minute points of metallic iron were found; in the granular portion none was found. The contrast between the two portions furnishes very satisfactory evidence of the fact of incongruent melting of synthetic fayalite. In their quantitative relations, however, the experiments are frankly disappointing, presumably because it is impossible to hold the fayalite in the liquid state an adequate length of time. We have postponed further investigation of this question until we find a container more nearly inert to molten fayalite than is silica glass.

Results Represented as a Binary System.

On account of the necessary presence of some Fe_2O_3 in the liquid phase our results require a diagram of more than two components for their accurate presentation. However, if the small amount of Fe_2O_3 is calculated as FeO , i.e., if all the iron of the mixtures is stated as FeO , all of our mixtures lie very close to the orthosilicate join, $\text{Ca}_2\text{SiO}_4\text{-Fe}_2\text{SiO}_4$, and there is much gain in convenience if the results are so presented. In Table III we give the composition of our mixtures

In the analysis of Rockport fayalite FeO was determined by the Pratt⁶ method modified. Iron and manganese were separated by a double basic acetate precipitation. Careful tests showed the absence of Al_2O_3 , CaO , and MgO . Total iron was also determined by direct titration with KMnO_4 after reduction with H_2S and gave an almost identical value with that from weighing Fe_2O_3 after the basic acetate separation and reprecipitation with NH_4OH . Water was determined by the Penfield tube method.

The results are given in Table II. The material proves to have a considerably higher manganese content than that in fayalite from the same locality analyzed by Penfield.

TABLE II.
Composition of fayalite from Rockport, Massachusetts.

SiO_2	29.75
Al_2O_3	none
Fe_2O_3	0.83
FeO	66.10
MnO	3.20
MgO	none
H_2O	0.19
	100.07

The fact that there is as much as 0.83 per cent Fe_2O_3 in this fayalite and that it still shows separation of iron on melting is of particular interest, for it furnishes indisputable evidence that a content of Fe_2O_3 at least somewhat greater than 0.83 per cent is necessary to carry fayalite out of the field of iron. The amount determined by analysis of the liquid, in synthetic material, is 2.25 per cent, as Table I shows.

We have also attempted to prepare pure fayalite synthetically and to demonstrate the separation of iron when it melts. To obtain most crystalline silicate compounds it is only necessary to prepare a liquid by melting together the ingredients in the proper proportion and then to bring about crystallization under appropriate conditions. With fayalite this method cannot be adopted because a liquid of the composition of fayalite does not exist, at least up to temperatures as high as the melting point of iron. The preparation of fayalite was, therefore, carried out by another method that is sometimes used for silicates, namely, by baking together its solid ingredients at a temperature below the melting point. In the case of fayalite,

* Op. cit.

temperature $1117 \pm 3^\circ$. The series of solid solutions also continues through the composition of the compound, CaFeSiO_4 , with the result that there is solid solution of the compound with the other end member, Ca_2SiO_4 , but in this case only a partial series of the type with-

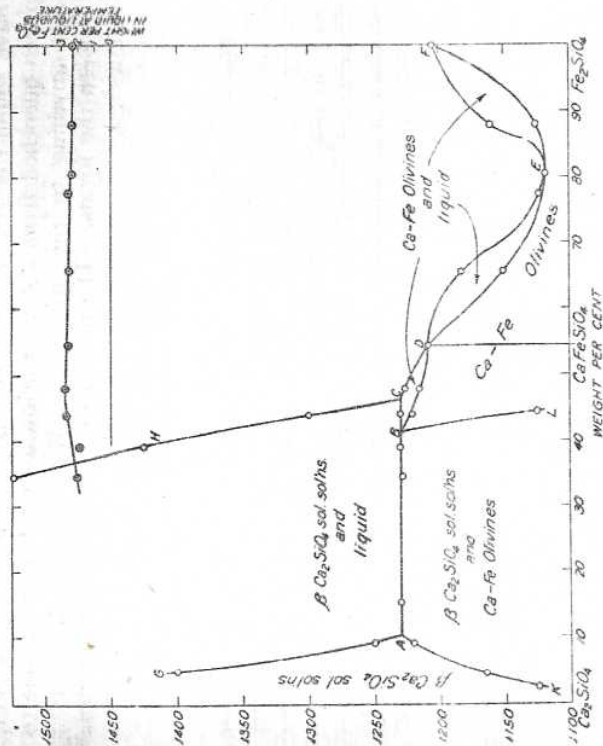


Fig. 2. Experimental results expressed in the form of a binary diagram by calculating all iron oxide as FeO . Upper right—Actual amounts of Fe_2O_3 in each liquid at liquidus temperature.

out a eutectic (Type IV, Roozeboom). The limit of solid solution in this series is at 41 per cent Fe_2SiO_4 (B, Fig. 2) at the point where solid solution is in equilibrium with liquid of composition 46 per cent Fe_2SiO_4 (C, Fig. 2) and Ca_2SiO_4 solid solution of composition 10 per cent Fe_2SiO_4 (A, Fig. 2) and the temperature is 1230° .

Calcium Orthosilicate and its Solid Solutions.

In earlier work of this Laboratory the compound, Ca_2SiO_4 , has been shown to occur in three forms termed α , β , and γ . The α -form is stable at temperatures from 1420° to its melt-

so calculated and the temperature of phase changes, and in Fig. 2 we give a "binary" diagram based upon this table.

TABLE III.
The data of Table I recalculated to a binary basis by expressing all iron oxide as FeO .

Composition		Temp. of beginning of melting	Temp. of completion of melting	Solid phases in equilibrium with liquid at liquidus temperature	Temperature
Fe_2SiO_4	Ca_2SiO_4				
100	0	1205 $\pm$ 2°	1205 $\pm$ 2°	Fayalite	1230 $\pm$ 2°
88	12	1125 $\pm$ 5°	1160 $\pm$ 2°	Ca-Fe olivine	1208 $\pm$ 2°
81	19	1117 $\pm$ 3°	1117 $\pm$ 3°	Ca-Fe olivine	1117 $\pm$ 3°
77	23	1120 $\pm$ 3°	1122 $\pm$ 3°	Ca-Fe olivine	1205 $\pm$ 2°
66	34	1150 $\pm$ 5°	1182 $\pm$ 2°	CaFeSiO ₄	
54.5	45.5	1208 $\pm$ 2°	1208 $\pm$ 2°	Ca-Fe olivine	
48	52	1215 $\pm$ 3°	1226 $\pm$ 2°	β -Ca ₂ SiO ₄	
44 ^a	56	1230 $\pm$ 2°	1300 $\pm$ 5°	β -Ca ₂ SiO ₄	
39	61	1228 $\pm$ 2°	1425 $\pm$ 5°	Ca ₂ SiO ₄ ^b	
34	66	1230 $\pm$ 2°	1525° (approx.)	α -Ca ₂ SiO ₄	
15 ^c	85	1230 $\pm$ 2°	2130 $\pm$ 20°	α -Ca ₂ SiO ₄	
0	100	1230 $\pm$ 20° ^d			

^a "Invariant points."

Composition of liquid
% Fe_2SiO_4 : 46, 54.5, 81, 100
Composition of solids
% Fe_2SiO_4 : 10, 41
Temperature of disappearance
1230 $\pm$ 2°, 1208 $\pm$ 2°, 1117 $\pm$ 3°, 1205 $\pm$ 2°

^a Additional determination on this mixture: Temperature of disappearance of Ca-Fe olivine and appearance of β -Ca₂SiO₄ = $1230 \pm 2^\circ$.

^b Probably α -form.

^c This mixture not made by complete fusion as were all others, but merely by mixing powders of Ca_2SiO_4 and mixture E (Fig. 2) and heating at various temperatures until temperature of beginning of melting was found.

^d Rankin and Wright, this Journal, 39, 7, 1915.

Discussion of Figure 2.

As shown in Fig. 2, Ca_2SiO_4 and Fe_2SiO_4 form the 1:1 compound, CaFeSiO_4 (54.5 per cent Fe_2SiO_4), analogous in composition to monticellite, CaMgSiO_4 . CaFeSiO_4 has a congruent melting point at $1208 \pm 2^\circ$. Crystals of the compound, CaFeSiO_4 , have frequently been found in slags but have not been recognized in nature. Between CaFeSiO_4 and fayalite there is a complete series of solid solutions of the type with a minimum melting point (Type III, Roozeboom).^a The minimum is at the composition 81 per cent Fe_2SiO_4 , and the

^a Z. phys. Chem., 30, 385-412, 1899.

ing point at 2130°, the β -form is stable between 675° and 1420° and the γ -form below 675°. The α - and β -forms have closely similar properties and their differentiation is not very satisfactory, but the α -form always shows a complicated twinning and the β -form only occasionally shows twinning.⁷ We have found the same contrast in charges containing Ca_2SiO_4 when quenched from above or below 1420°, but are unable to add anything to the information regarding the distinction between the forms. The indices of these forms are given as follows⁸:

	γ	β	α
α -form	1.737	1.720	1.715
β -form	1.735		1.717

The α - and β -forms are obtainable only with difficulty on account of their strong tendency to invert to the γ -form on cooling, but with very rapid cooling and especially when they occur as small crystals embedded in another phase, the higher-temperature forms may sometimes be obtained at room temperature and examined microscopically. Only occasionally did we obtain grains of the α - or β -forms in our iron-bearing mixtures. In these cases it was not possible to measure the properties of the crystals with great accuracy but there was no obvious change in the properties of these forms when they grew in liquids containing some 45 per cent Fe_2SiO_4 , except perhaps that the birefringence of the β -form seemed to be somewhat lowered. On the other hand, the γ -form produced by inversion of such grains of β - Ca_2SiO_4 has refractive indices notably higher than those of pure γ - Ca_2SiO_4 . From this fact we concluded that the β -form must have contained iron in solid solution before its inversion, even though its properties were not obviously changed,⁹ and we, therefore, investigated the limit of the solid solutions. To do this we mixed pure Ca_2SiO_4 as a very fine powder with varying amounts of material of the composition of the minimum E (Fig. 2) and held the mixture in iron foil at 1220°, i.e., about 10° below the horizontal ABC. At this temperature the material of composition E forms a liquid which soaks into the Ca_2SiO_4 and forms a solid solution with it. It was found that all mixtures up to a concentration corresponding with the point A

⁷ Day and Shepherd, this Journal, 22, 280-284, 1906.

⁸ Rankin and Wright, this Journal, 39, 7-8, 1915.

⁹ That β - Ca_2SiO_4 should suffer no great change of mean refractive index through solid solution of CaFeSiO_4 is not surprising since their mean refractive indices are not far apart, but a very different condition prevails in the case of γ - Ca_2SiO_4 .

(10 per cent Fe_2SiO_4) were thus converted into homogeneous solid solutions. Beyond the point A (more than 10 per cent Fe_2SiO_4) an excess phase appeared and this corresponded in properties with the saturated Ca-Fe olivine (point B). In this manner the limit of solid solution of CaFeSiO_4 β - Ca_2SiO_4 was determined as 10 per cent Fe_2SiO_4 (A).

It should be pointed out that, although the method described gave the limit of β - Ca_2SiO_4 solid solutions, the material examined under the microscope had in all cases inverted partially and in some cases completely to the γ -form. There is a progressive change in the readiness of inversion depending upon the composition. Thus pure β - Ca_2SiO_4 can be cooled rapidly enough so that a fair proportion remains in that form. With 5 per cent Fe_2SiO_4 in solid solution the cooling must be very rapid to retain even a small proportion in the β -form, and with 10 per cent Fe_2SiO_4 in solid solution we were unable to prevent complete inversion to the γ -form. It was upon such γ - Ca_2SiO_4 solid solutions of known composition that the change of refractive indices of this form with composition was determined.

Slopes of the Curves AK and BL.

Obviously, the method used to determine the point A can be used to determine other points on the curve AK and likewise points on the curve BL. The lower limit of temperature to which the method may be extended is, however, that of the point E (1117°), for to attain equilibrium there must be melting of the mixture E to a liquid in order that this liquid may soak into the Ca_2SiO_4 and give the appropriate solid solution. At a temperature at which no liquid forms, equilibrium proceeds too slowly to permit its determination. By determining the composition at which Ca-Fe olivine appears as an excess phase at 1165° and at 1125° the curve AK was fixed. It will be noted that at lower temperatures there is a very marked decrease in the amount of the iron compound taken into solid solution. At 1125° the saturated solid solution contains only about 2 per cent Fe_2SiO_4 . By working in a similar manner with compositions in the neighborhood of the curve BL and determining the composition in which a small amount of Ca_2SiO_4 appears together with the Ca-Fe olivine, the limit of concentration of the Ca-Fe olivine at various temperatures was determined and thus the curve BL located. It will be noted that the curve BL slopes to the right downward.

Calcium Orthosilicate Solid Solutions in the γ -form.

It has been stated above that the β - Ca_2SiO_4 solid solutions inverted either completely or partially to the γ -form on cooling to room temperature. The result was that the material examined under the microscope appeared in the γ -form with not more than an occasional remnant of β -form. The γ -solid solutions thus formed by inversion have indices notably higher than pure γ - Ca_2SiO_4 . It would, of course, be possible for β -solid solutions to invert to the γ -form with separation of Ca-Fe olivine in so fine a form that the microscope could not detect it, in which case the raised refractive indices would be merely aggregate refractive indices. That this is not true and that there is real solid solution in the γ -form is shown by the fact that the X-ray powder pattern shows a change of spacing of the lines (see Fig. 6). We have not found it possible to determine the limit of the series of solid solutions in the γ -form. When one prepares them by inversion from the β -form one cannot, of course, go beyond the limit of solution in the β -form, viz., 10 per cent Fe_2SiO_4 . There is no reason for believing that solution in the γ -form stops at this point, but attempts to prepare γ -solid solutions richer in iron by baking together solid ingredients at temperatures from 650-750° gave only nondescript products even with the aid of a flux.¹⁰ The attempts were made particularly because a recent investigation by Greer suggests that γ - Ca_2SiO_4 forms a complete series of solid solutions with Mn_2SiO_4 ,¹¹ and it might be expected that similar relations would hold with Fe_2SiO_4 . It must be admitted that the change of refractive indices of the γ -form with composition is such that the extrapolated curves might be connected up with the extrapolated curves for the Ca-Fe olivines though this would necessitate strong curvature of the missing portion of the curves and the determined portions seem to be nearly, if not quite, straight (see Fig. 5). On the other hand, some of the thermal results indicate that there cannot be complete solid solution with the γ -form, and these are the results which fixed the curve BL (Fig. 2). The determined slope of the curve BL necessitates that the completed diagram showing the relations at lower temperatures should have the general form shown in Fig. 3. There is no means of predicting the actual length of ST, representing the

¹⁰ It was necessary, of course, to carry on this work in evacuated, sealed tubes to prevent oxidation of iron to the ferric state.

¹¹ Greer, W. L. C., *Am. Mineral.* 17, 135-142, 1932.

magnitude of the break in the solid solution series, but complete solid solution could occur only if S and T coincide, when the curves BLT and SV become a single curve as shown in Fig. 4 (VR and RA likewise become a single curve)

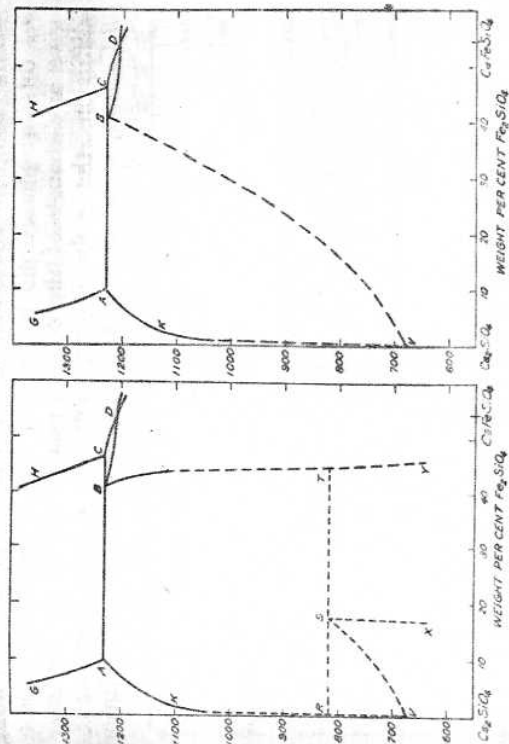


Fig. 3. Hypothetical completion of the equilibrium diagram at lower temperatures in the general form necessitated by the determined slope of BL.
Area RABT = β - Ca_2SiO_4 solid solutions and Ca-Fe olivines.
Area VSR = β - Ca_2SiO_4 and γ - Ca_2SiO_4 solid solutions.
Area VSX = γ - Ca_2SiO_4 solid solutions.
Area XSTY = γ - Ca_2SiO_4 solid solutions and Ca-Fe olivines.

Fig. 4. Hypothetical completion of the equilibrium diagram in the general form necessary if γ - Ca_2SiO_4 formed a complete series of solid solutions with CaFeSiO_4 .

Area VAB = β - Ca_2SiO_4 solid solutions and γ - Ca_2SiO_4 solid solutions.

Area bounded upward and to the left by

VBD = γ - Ca_2SiO_4 solid solutions or Ca-Fe olivines, which become synonymous terms.

Obviously, this type of diagram can obtain only when the curve BV slopes downward to the left throughout its length, whereas our determined portion BL has the opposite slope. These considerations have led us to conclude that the diagram as completed at lower temperatures must have the general form of Fig. 3, presenting a break in the solid solution series.

It should, perhaps, be emphasized that the difficulty of determining the equilibrium conditions at these lower temperatures is due to the fact that one cannot cool any of these mixtures

from the high temperatures at which they are completely molten without obtaining crystallization of $\beta\text{-Ca}_2\text{SiO}_4$ solid solutions on passing through the region GACH (Fig. 3) and without crystallization of Ca-Fe olivine in the region RABT, giving complete consolidation. Even several days' heating of this two-phase solid material at low temperatures fails to give any change of the composition of the phases by interdiffusion, the only change obtained being inversion of $\beta\text{-Ca}_2\text{SiO}_4$ solid solution as such to $\gamma\text{-Ca}_2\text{SiO}_4$ solid solution

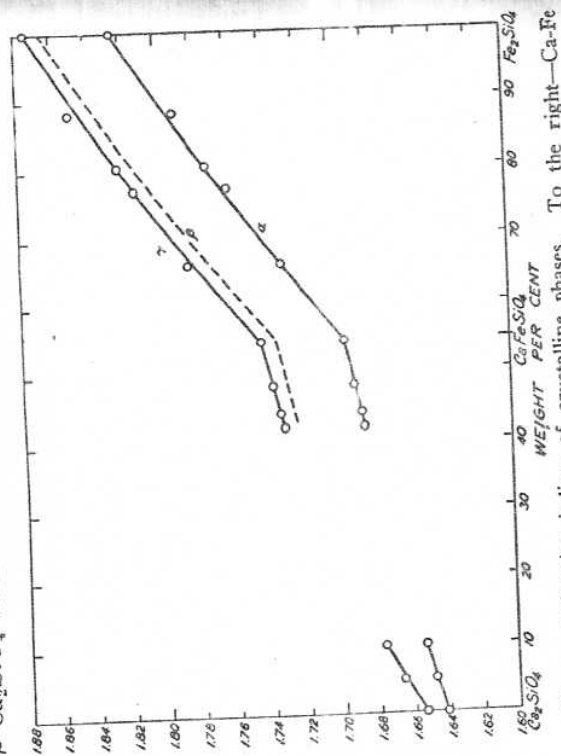


Fig. 5. Refractive indices of crystalline phases. To the left— $\gamma\text{-Ca}_2\text{SiO}_4$ solid solutions, olivines. To the right— $\gamma\text{-Ca}_2\text{SiO}_4$ solid solutions, olivines.

of the same composition, the Ca-Fe olivine remaining unchanged. It is difficult to see how conditions can be greatly different in the corresponding Mn system. The temperatures must rise to the same high values at the Ca_2SiO_4 end of the diagram. Cooling through the region where $\beta\text{-Ca}_2\text{SiO}_4$ is stable must likewise result in the separation of that phase so that Greer's results indicating complete solid solution in the γ -phase, obtained by the simple method of cooling a liquid, would seem to necessitate the conclusion that in the Mn system there is complete solid solution in the β -phase as well and that γ -form solutions formed at high temperatures inverted to the γ -form without change of composition.

OPTICAL PROPERTIES OF THE SOLID SOLUTIONS.

The Ca-Fe Olivines.

The refractive indices of the crystalline phases were measured in immersion liquids in sodium light. For the Ca-Fe olivines and for $\gamma\text{-Ca}_2\text{SiO}_4$ solid solutions the results are given in Table IV and plotted in Fig. 5, only γ and α being determined for each composition. The exact character of the

TABLE IV.

Refractive indices of crystalline phases.

Ca-Fe Olivines		
Composition		
Fe ₂ SiO ₄	Ca ₂ SiO ₄	
100	0	γ 1.874 ± .003
88	12	α 1.824 ± .003
81	19	1.790*
77	23	1.772
66	34	1.760*
54.5	45.5	1.730*
48	52	1.696
44	56	1.690
41	59	1.737
		1.686
		1.684
		1.653 ± .002
10	90	1.647
5	95	1.642
..	100	1.634

* Slight zoning of crystals present in these compositions.

material used requires some discussion. The crystalline material, in the case of the Ca-Fe olivines, was made merely by permitting a liquid to crystallize upon cooling. These liquids always contained some Fe_2O_3 , but upon crystallization with moderately slow cooling, this Fe_2O_3 is not uniformly distributed in the crystals for it can be observed that the core of the crystals is nearly colorless, whereas the rims are notably colored. There is thus a concentration of Fe_2O_3 in the portions of later crystallization. In powder under the microscope more highly colored grains are found to have higher refractive indices than the colorless grains, a condition that is to be expected from their content of Fe_2O_3 . In measuring the refractive indices these colored grains were avoided and we thus believe that our results refer to the almost purely ferrous composition.

In some of the mixtures there is zoning of another kind, namely, of the ordinary type that could be predicted from inspection of the equilibrium diagram (Fig. 2). Indeed, it was found impossible to crystallize those mixtures for which the solidus and liquidus temperatures are far apart (as for example the mixture with 88 per cent Fe_2SiO_4) without obtaining some effect of that kind unless the cooling was made so rapid that only feathery crystals, not well adapted to index measurements, were obtained. It was necessary in dealing with such mixtures to strike a compromise between a slow rate of cooling which gave the former effect and a rapid rate giving the latter. Fortunately, there are three compositions in which zoning of that kind is impossible, namely, that of the compound, CaFeSiO_4 , that of the compound, Fe_2SiO_4 , and that of the minimum melting temperature (E, Fig. 2). These can be cooled as slowly as one may wish and thus crystalline material obtained that is entirely suitable for accurate measurement of indices. In drawing the curves of variation of refractive indices with composition (Fig. 5) greatest weight has been given to these three compositions, in fact the curves were drawn through the points determined for them. It will be noted that for other mixtures there is a tendency for the points representing γ to lie somewhat above the curve and for those representing α to lie somewhat below the curve. The nature of the deviation is thus always of the kind that is to be expected in material that is slightly zoned and in which maximum and minimum refractive indices have been measured statistically. The magnitude of the deviation is always small and does not destroy the plain evidence of continuous variation with composition.

The birefringence of the Ca-Fe olivines changes only a small amount from end to end of the series, $\gamma - \alpha$ being 0.050 at fayalite, 0.47 at CaFeSiO_4 , and 0.045 at the lime-rich extreme which has 41 per cent Fe_2SiO_4 . All are optically negative and 2V is sensibly constant throughout at $50 \pm 2^\circ$. An interesting feature of the series is the very marked change of slope of the curves of refractive indices at the composition of the compound, CaFeSiO_4 .

The values we have obtained for fayalite, $\gamma = 1.874$, $\alpha = 1.824$, are practically the same as those found by Penfield for fayalite from Rockport, Massachusetts. The material measured by Penfield analyzed nearly pure fayalite. Considerably higher values than these ($\gamma = 1.886$, $\alpha = 1.835$) based upon

measurements by Busz¹² are sometimes given for fayalite. The crystals came from a slag and in the light of our measurements there is reason to believe that they must have contained some material in solid solution which raised their indices above those for pure fayalite.

The $\gamma\text{-Ca}_2\text{SiO}_4$ Solid Solutions.

Solid solution of Fe_2SiO_4 in $\gamma\text{-Ca}_2\text{SiO}_4$ up to 10 per cent Fe_2SiO_4 , which limits the series as prepared by us, induces a rapid rise of refractive indices, the measured values being given in Table IV and plotted in Fig. 5. For reasons already stated in the discussion of the equilibrium studies the series is to be regarded as extending at least somewhat farther. On account of intricate twinning we were unable to measure any of the optical properties except γ and α with any degree of satisfaction. This twinning shows on the sections giving maximum birefringence as three sets of lamellae cutting each other at approximately 60° . Occasionally the number of lamellae is reduced so that the grain is divided into six sextants with only an extra lamella or two, giving trillings analogous to those seen in aragonite, and in such sections the extinction is symmetrical. The more familiar orthosilicates, the rock-forming Mg-Fe olivines, occasionally show twinning on the dome (011) which gives trillings. Possibly we have in the present $\gamma\text{-Ca}_2\text{SiO}_4$ solid solutions a twinning after a similar law and the apparently complex twinning is not necessarily to be taken as indicating monoclinic symmetry. The $\gamma\text{-Ca}_2\text{SiO}_4$ series is possibly orthorhombic.

It should be noted that in the Ca-Fe olivines of our preparations no twinning of any kind was observed.

THE X-RAY POWDER PATTERNS OF THE CRYSTALLINE PHASES.

Together with the thermal and optical investigation of the system, $\text{Ca}_2\text{SiO}_4\text{-Fe}_2\text{SiO}_4$, an X-ray study by the "powder method" was also made. The diffraction patterns obtained from a number of compositions in this system are reproduced in Fig. 6. They were made with a General Electric diffraction apparatus using molybdenum K_α radiation. Their examination will readily show that starting from pure Fe_2SiO_4 (fayalite), and increasing by equal steps the Ca_2SiO_4 content to over

¹² Busz, K., and Rüsberg, F. W., Central. Min., p. 625, 1913.

60 per cent by weight, a continuous displacement of the diffraction lines takes place without, however, changing the general characteristics of the diffraction pattern. The interpretation of this as existence of a series of solid solutions in this portion of the system is obvious, and the observed dis-

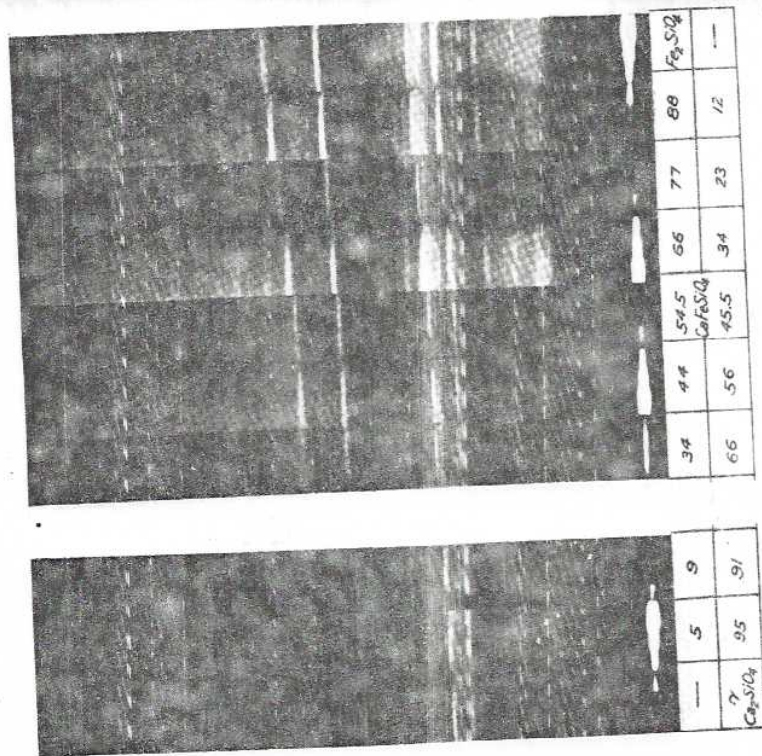


Fig. 6. X-ray diffraction patterns of γ -Ca₂SiO₄ solid solutions and of Ca-Fe olivines.

placement of lines in the spectrograms results from the expansion of the crystal lattice which would be expected from substitution of the Ca²⁺ for the smaller Fe²⁺ (the ionic radii are 1.06 and 0.83 respectively). It should be noticed at the same time that the displacements are quite large up to the composition with 45.5 per cent Ca₂SiO₄ (CaFeSiO₄), and that with a further increase in the calcium content a relatively

much smaller additional displacement takes place. It will be recalled also that the rate of change of refractive indices becomes much less at the composition of the compound (see Fig. 5). The preparation containing 66 per cent Ca₂SiO₄ lies beyond the limit of the Ca-Fe olivines and is found, when examined optically, to contain some γ -Ca₂SiO₄. The amount of the second phase is, however, not great enough to show in the powder spectrogram on account of the large number of diffraction lines from the principal constituent. Beyond the statement of the existence of a series of solid solutions to over 60 per cent content of Ca₂SiO₄, the X-ray data do not permit further direct conclusions. They support, however, the conclusion drawn from the thermal study which gave proof of the existence of the compound, CaFeSiO₄, in that they show that a different amount of displacement of the diffraction lines takes place on each side approaching the composition corresponding to this compound. The combined evidence proves complete miscibility from Fe₂SiO₄ to CaFeSiO₄ as well as some further miscibility of Ca₂SiO₄ and CaFeSiO₄; also that no unmixing of these solid solutions has taken place on cooling to room temperature.

It was mentioned above that the preparation containing 66 per cent of Ca₂SiO₄ was found to contain two phases when examined microscopically. Beyond this composition no homogeneous products were obtained until the Ca₂SiO₄ content was increased to about 90 per cent. The X-ray diffraction patterns obtained from melts containing 91, 95, and 100 per cent Ca₂SiO₄ were identical except for a small offset of the lines (see Fig. 6). The displacement, which represents a contraction of the lattice relative to pure Ca₂SiO₄, and is due to substitution of the larger Ca²⁺ by Fe²⁺, proves that solid solutions exist also in this portion of the system.

It has already been shown in another part of the paper that β -Ca₂SiO₄ takes up a maximum of 10 per cent Fe₂SiO₄ in solid solution and that, since the γ -Ca₂SiO₄ was obtained only by inversion from the β -form, the limit of concentration of Fe₂SiO₄ in the γ -solid solutions obtained by us was necessarily 10 per cent. It was further shown that solid solution of Fe₂SiO₄ in γ -Ca₂SiO₄ must extend at least a little farther, but complete miscibility extending over to the Ca-Fe olivine series was excluded by the thermal results. The X-ray diffraction evidence taken by itself tends to confirm the existence of an hiatus. The diffraction patterns of the two series

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exhibit differences which suggest a distinct difference in their symmetries.

In this connection it is of interest to recall that in the study of the system, $\text{Ca}_2\text{SiO}_4\text{-Mn}_2\text{SiO}_4$, Greer¹³ stated that in this case there is an unbroken series of solid solutions. He mentions some difficulties in preparing members of this series containing 50, 60, and 70 per cent Ca_2SiO_4 , which are apparently overcome by annealing at 1100°. What this heat treatment does is not clear, and his presentation does not permit independent judgment.

Two other studies of interest here are the crystal structure determinations of forsterite¹⁴ and monticellite.¹⁵ These minerals were found to have identical structures. This is possible because the eight magnesium atoms in the unit cell of forsterite do not occupy equivalent positions, but are divided into two groups of four each, and it is possible to replace one group of magnesium atoms by calcium without a change in symmetry. The close similarity of forsterite and monticellite on one hand, with that of fayalite and CaFeSiO_4 on the other, would suggest that a similar structural relation exists in the latter case. This would furnish a plausible explanation for the existence of the unbroken series of solid solutions which has actually been found between Fe_2SiO_4 and CaFeSiO_4 . The explanation, however, becomes less convincing in view of the fact that there is much doubt regarding the existence of continuous solid solutions between forsterite and monticellite; firstly, because the available analyses of the respective minerals fail to show any members of intermediate composition, and secondly, because the investigation of the system, CaO-MgO, SiO_2 , by Ferguson and Merwin¹⁶ indicated a miscibility of about ten per cent (their experiments, however, give only the lower limit). The elucidation of these interesting questions requires further investigation.

PETROLOGIC APPLICATIONS.

It is rather surprising that Ca-Fe olivines of the series we have prepared are not found in nature. The most likely manner of occurrence would be in contact metamorphic deposits involving metasomatism. Perhaps the oxides concerned more

readily form garnet under such conditions especially if much ferric iron is present. Nevertheless, it does seem probable that these olivines should occur. The optical properties are not strikingly different from those of the more familiar Mg-Fe olivines and mere observation of relief and interference colors in thin section might not serve to distinguish them. With careful study and especially with measurement of refractive indices the distinction is readily made, for in the Ca-Fe series the birefringence is much greater for a given mean index of refraction than in the Mg-Fe series.

The finding that CaFeSiO_4 forms a complete series of solid solutions with fayalite, which in turn almost certainly forms a complete series with forsterite, renders it rather surprising that the ordinary rock-forming Mg-Fe olivines do not contain more CaO. The content of CaO in these olivines is ordinarily very low, such an amount as 2 per cent being apparently exceptional, though as much as 5 per cent has been noted in the olivine of a rock type of alkaline affinities from Cape Verde Islands. The significance of the low lime content of most rock-forming olivines may become apparent after a thermal study of the system involving all three of these orthosilicates.

The occurrence of the natural mineral larnite is very remarkable. Although the distinction between α - and β - Ca_2SiO_4 is not as clear as could be desired, the properties of larnite agree best with those of the α -form as Tilley has pointed out.¹⁷ This form, in its pure state, is stable only above 1420° and the other forms are quite unknown as natural minerals. We have here a very unusual state of affairs. Ordinarily, it is the low-temperature forms of silicate compounds that are found in nature. Thus, of the two forms of the metasilicate of lime, wollastonite and pseudowollastonite, only wollastonite is known in nature and of the two forms nephelinite and carnegite only nephelinite is found. Or, if more than one form occurs naturally, the low-temperature form is the common one, the others relatively rare, as for example in quartz, tridymite, and cristobalite.

Another remarkable feature of larnite is its survival. The α -form of Ca_2SiO_4 and the β -form as well are extremely difficult to prepare in any quantity on account of their marked tendency to change to the γ -form. Yet in larnite we have one

¹⁷ Op. cit., p. 81.

¹³ Greer, W. L. C., *Am. Mineral.*, 17, 135, 1932.

¹⁴ Bragg, W. L., and Brown, G. B., *Z. Krist.*, 63, 538, 1926.

¹⁵ Brown, G. B., and West, J., *Z. Krist.*, 66, 154, 1927.

¹⁶ Ferguson, J. B., and Merwin, H. E., *this Journal*, 48, 81, 1919.

of these forms, probably α , made in quantity in nature and persisting through the period of very slow cooling that must have prevailed after its formation. In our study of Ca_2SiO_4 containing Fe_2SiO_4 in solid solution we found that the tendency to invert increases with the content of Fe_2SiO_4 . It is probable that natural larnite contains some material in solid solution that has the opposite effect. Tilley's analysis shows 1.12 per cent Al_2O_3 and 0.69 per cent MgO which he says "are probably represented in finely divided spinel,"¹⁸ but the spinel, separately analyzed, shows much Fe_2O_3 , whereas the larnite analysis gives only a trace. It seems doubtful, therefore, that the content of extraneous oxides is entirely accounted for by inclusions. There is probably some solid solution and possibly the material in solid solution decreases the inversion tendency of the larnite. It is certain, at least, that the natural larnite, which we have heated under varying conditions, has a distinctly smaller tendency to invert than pure Ca_2SiO_4 . Nevertheless, inversion can be induced, though with varying ease in different specimens, and there is still the problem of the survival of the larnite in nature. Perhaps the pressure to which larnite was subjected during cooling was sufficient to prevent the change to the γ -form, which is accompanied by a 10 per cent increase of volume. This change of volume is particularly large and of the opposite sign to that usually observed with silicates, the density of the low-temperature form being ordinarily greater than that of the high-temperature form. High pressure should, therefore, induce a marked lowering of the inversion temperature of larnite and it is possible that, during its formation the pressure may have been such that no excessively high temperature was necessary in order that the larnite should have formed within its stability range. The temperature may not have approached the value 1420° , which is the minimum temperature of stable existence of pure $\alpha\text{-Ca}_2\text{SiO}_4$ at ordinary pressure.

APPLICATIONS TO SLAGS.

The slags of widest use in non-ferrous metallurgy are those made up principally of CaO , FeO , and SiO_2 and the monosilicates (orthosilicates) are not uncommonly employed. Since fayalite contains about 30 per cent SiO_2 , and Ca_2SiO_4 about 35 per cent, all intermediate compositions contain intermediate amounts of SiO_2 so that all lime-iron slags with a little more

than 30 per cent SiO_2 lie close to the orthosilicate join. More siliceous slags, approaching the metasilicates, are usually preferred in the copper blast furnace because the loss of copper in the slag is less than with the orthosilicates. In the copper converter, however, orthosilicate slags are used, a composition with a very high iron content approaching fayalite being common. It is to be noted that fayalite forms a liquid at 1205° , a temperature considerably lower than the value commonly stated in metallurgical literature ($1250\text{-}1270^\circ$).

In lead smelting a wide range of orthosilicate slags have been used. Our results do not, of course, enable us to predict anything about metal losses and other factors of great importance in the practical working of slags. We can say only that the orthosilicate slag of lowest melting temperature is that with 81 per cent Fe_2SiO_4 . Though it was not part of our work to measure fluidity, nevertheless, impressions were gained during the handling of the mixtures and it appeared that at any given temperature the fluidity increased with the iron content so that this minimum-melting mixture is also a very fluid slag.

Other investigators have made measurements of the melting temperatures of mixtures corresponding with the orthosilicate series here studied. Among these we shall mention only Loo and Kern, who made cones of several orthosilicate mixtures and observed their deformation temperatures.¹⁹ Fig. 7 permits comparison of their results with ours, and shows that there is good general agreement. We may note, however, that their mixture No. 31 (see Fig. 7) corresponding with 29 per cent Fe_2SiO_4 71 per cent Ca_2SiO_4 , for which they give a "fusion temperature" of 1250° , could have been no more than about 55 per cent liquid at that temperature and a rise of temperature of 400° is necessary before complete fusion occurs. This is a striking example of the inadequacy of observations of softening and flowage as a means of determining fusion temperatures of mixtures. Our results show that in orthosilicate slags the steeply rising fusion surface of Ca_2SiO_4 is encountered at a point corresponding with about 35 per cent CaO . With a greater proportion of CaO melting may appear to take place at 1230° , but it is only partial melting and if one attempted to use an orthosilicate slag with more than 35 per cent CaO at a temperature in the neighborhood of 1230° difficulties would certainly be encountered as a result

¹⁹ Trans. Am. Inst. Mining and Metallurg. Eng., 76, 503-505, 1928.

¹⁸ Op. cit., p. 81.

of segregation of crystals of Ca_2SiO_4 . Among the 25 mixtures that Loo and Kern used in actual tests of their efficacy as practical slags this one was not tried. Of the four mixtures selected by them as "the best slags to produce in the smelting of roasted cassiterite concentrates" two were the orthosilicate mixtures, Nos. 14 and 15 (Fig. 7). It will be noted that

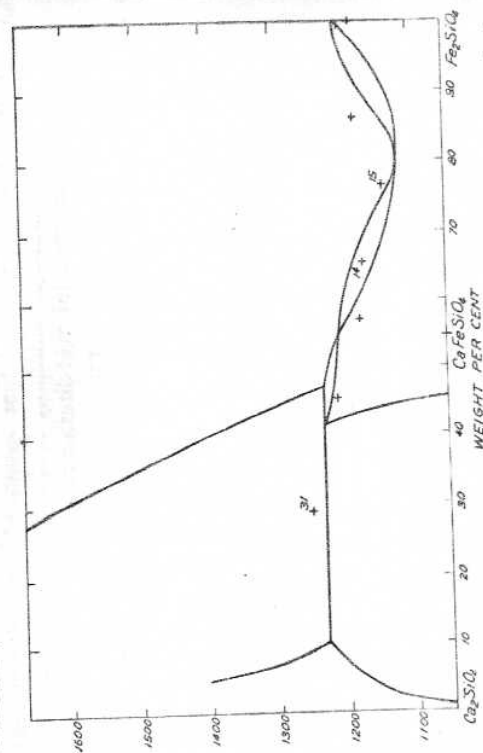


Fig. 7. Comparison diagram showing in full curves the results of the present investigation and by crosses "Fusion Points" determined by Loo and Kern (op. cit., p. 504), whose "Mixture No." is given for three of these points.

these, and especially the latter, are not far from the composition of the minimum-melting mixture.

In the basic open-hearth process for steel the slag as first melted may be not far from orthosilicate composition so that our determinations have some application to that process in its early stages. At later stages the slag ordinarily acquires large amounts of phosphorus, the elimination of which is an important feature of the process, and the slag then departs notably in composition, and, therefore, in its thermal relations, from any purely siliceous slag.

SUMMARY.

The system, $\text{Ca}_2\text{SiO}_4\text{-Fe}_2\text{SiO}_4$, has been studied thermally by using iron crucibles as containers and an atmosphere of nitrogen to prevent oxidation. Even in the presence of metal-

lic iron not all of the iron oxide of these mixtures is reduced to the ferrous state, but an equilibrium is established when there is still a small amount of Fe_2O_3 . Accurate presentation of the results requires a ternary diagram in order to show the presence of this small amount of Fe_2O_3 . This is done in Fig. 1. A great gain in the utility of the results is made if the small amount of Fe_2O_3 is calculated as FeO and thus all the iron oxide expressed as FeO . We thus obtain a "binary" diagram from which temperatures and compositions are easily read. Such a diagram is shown in Fig. 2. The diagram shows that there is one intermediate compound, viz., CaFeSiO_4 , which melts at $1208 \pm 2^\circ$ and forms a complete series of solid solutions with fayalite which melts at $1205 \pm 2^\circ$. The solid solution series is of the type with a minimum (Type III, Roozeboom) and the minimum is at $1117 \pm 3^\circ$ with a composition 81 per cent Fe_2SiO_4 .

In addition CaFeSiO_4 and $\beta\text{-Ca}_2\text{SiO}_4$ form a series of solid solutions of the type with an hiatus and with all melting temperatures intermediate between those of the end members (Type IV, Roozeboom).

The existence of these solid solutions is confirmed by optical evidence, which is presented in Fig. 5, and by X-ray evidence, presented in Fig. 6.

The results are discussed in connection with certain problems of the petrologist and of the metallurgist.

ACKNOWLEDGMENTS.

We are indebted to Dr. H. E. Merwin for aid in the use of high-refracting liquids of his devising, and to Dr. J. W. Greig for the identification of metallic iron in the fayalite melt.

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SiO₂-content was too low to form the normal standard minerals; in these cases the deficit of Q is given, instead of the calculated amount of olivine or spinels.

The first group of 7 analyses (without much Fe-cordierite or sillimanite in the kata standard norm) corresponds to rocks with only subordinate quantities of special minerals. If the content of Fe-cordierite and sillimanite is a little higher, garnet (almandine) is more easily formed in the mesofacies (the next 2 analyses). Higher content of Fe-cordierite and (or) sillimanite is found in the last 3 analyses which correspond to meso rocks rich in staurolite.

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FAYALITE AT ROCKPORT, MASSACHUSETTS*

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ABSTRACT

The facts are collected concerning the place and mode of occurrence of fayalite at four localities on Cape Ann, Massachusetts, with a description of one of them which yielded the largest and best-developed crystals. All of these occurrences are in pegmatite of a granitic nature but evidence is presented which seems to show that all are xenoliths, genetically related to fayalite-bearing nordmarkite granite, older than and intruded by the dominant Rockport granite.

Fayalite is not a mineral of common occurrence. It is true that it has been found as a constituent of certain granites and is well known to be abundantly formed in the slags of iron smelters. But individual crystals of natural occurrence are very rare. Boulders of an iron silicate found on a beach on the island of Fayal in the Azores were supposed to have come from the local volcanic rocks and the name *fayalite* was based on their description. However, further study showed that in all probability the boulders were in reality lumps of slag, carried as ballast in some ship and left ashore there. Nevertheless the name has persisted. Minute crystals of great beauty have been found in the cavities of rhyolites and from them the crystal form has been well established.

These facts are stated in order to bring out the very unusual character of the four occurrences of fayalite known in the granite of Rockport on Cape Ann, Massachusetts, to the description of which the following pages are devoted.

The first to find fayalite at Rockport was Mr. J. H. Sears (1905), the Curator of the Peabody Museum at Salem. In 1890 he sent specimens to Professor Penfield at Yale University, reporting that they had been found in the granite of the large Rockport Granite Quarry at a depth of sixty feet. They were parts of a lenticular shell of varying thickness, from twelve to sixteen inches in diameter, filled on the inside with loose earthy material and enveloped by a layer of magnetite about one inch thick. This was in the granite but adjacent to a pegmatite boss or vein. There is a piece of this find in the Harvard Collection which shows a section of a large crystal like those to be described later. On fresh fracture the color is greenish black with resinous luster and two cleavages are shown. Penfield (1896) showed that these cleavages were very perfect parallel to the basal pinacoid; less perfect parallel to the branchipinacoid, thus correcting the previous description of this property. He made an analysis, finding it to be a very pure silicate of iron; and he determined for the first

* Contribution from the Department of Mineralogy and Petrography, Harvard University, No. 319.

time its optical constants. Much later (Bowen and Schairer, 1932) it was found that the melting temperature of this fayalite was $1205^{\circ} \pm 2^{\circ}\text{C}$, exactly the same, within the limits of error, as that of pure synthetic ferrous silicate.

Fayalite was next reported from Rockport by Dr. Charles H. Warren, (1903) then of the Massachusetts Institute of Technology, who described a second discovery in the same quarry. This mass was found by two of his students and he did not see it in place. It was large, some 250 pounds having been obtained, and, as this was sold to dealers, the mineral was widely distributed in collections. The specimens were sections of large crystals, tapering from a thickness of ten inches down to a thin edge. There was a small amount of black lepidomelane mica on the surfaces of the crystals as well as minute zircon crystals. The main purpose of this paper was to describe a reaction zone developed where the fayalite was in contact with quartz; the zone consisted of a thin layer of a finely fibrous ferroan anthophyllite mixed with granular magnetite. It is now known (Bowen and Schairer, 1935) that this amphibole is grunerite, it being monoclinic rather than orthorhombic as Warren thought.

We next hear of fayalite at Rockport in 1908 and, as the writer had a hand in obtaining these specimens which are undoubtedly the finest found anywhere of this mineral and have never been adequately described, the facts may well be recorded in detail.

In 1908 Dr. Dale of the United States Geological Survey wrote on the Commercial Granites of Massachusetts. In describing the Rockport Granite he states that in the Babson Farm Quarry, which had been opened not long before on the extreme northeast point of Cape Ann, there had been found a "knot" or pegmatite of unusual character which contained, among other more usual minerals, "a mass six by two inches of yellowish brown color, determined by Mr. Johannsen as bronzite." As this mineral seemed wholly out of place in the paragenesis of granite pegmatite I at once suspected that another find of fayalite had come to light and took an early occasion to visit the quarry with a party of students.

We found the quarry itself well worth the journey. It had been opened by a contractor to supply rectangular granite blocks weighing thirty or forty tons each for the construction of a subsurface breakwater to protect the harbor of Rockport. The operation was on a gigantic scale, and the great blocks in process of being broken out of the quarry with wedges or hoisted out onto the surface for transfer by rail and scow were very impressive. The granite was coarse and wonderfully homogeneous with almost no sign of the dikes or pegmatite segregations so common in other Cape Ann quarries. By the side of the railway track stood the block which Dale had described, condemned because of its poor quality and

awaiting removal to the dump of waste rock. It measured about eight feet by four by two and a half, and was composed in large part of coarse pegmatite in sharp contact with the normal granite. The striking feature of this contact was its cross-cutting nature; the structure of the pegmatite bore no relation to it. The evidence was conclusive that the pegmatite was but a part of a larger mass which had been disrupted by the intrusion of the granite and that this angular block was one of the fragments, floated off as a xenolith in the granite. Search of the large waste dump and examination of the vast exposed faces of the quarry walls revealed no other like material. So this lone block, left by chance after being brought to the surface, to be seen by chance by a visiting geologist on a timely visit, was all there was found of this pegmatite.

The "bronzite," or rather fayalite as it proved to be, showed on two surfaces of the quarry block but getting it out looked like a hopeless task. However, two of the party, George M. Flint and A. C. Gosse, returned the next day armed with drills, hammers and plug-and-feather wedges. They obtained permission to break into the discarded stone, and ultimately brought back to Cambridge the larger part, perhaps a hundred pounds, of the pegmatite inclusion.

The chief constituent and latest mineral to form was quartz in large anhedral, glassy but of a peculiar hazy opalescent blue color; masses of several pounds weight were obtained. The feldspar was white microcline characterized by an unusual curvature both of platy masses and of individual crystals where it was embedded in quartz. In the feldspar were warped and cracked plates of ilmenite up to five inches in diameter but less than an eighth of an inch in thickness. There were also quite large prismatic crystals of a green pyroxene which proved to be hedenbergite. And lastly there was the fayalite, the large crystals of which were segregated in a single mass, lying against one another like sardines packed in a can. The principal specimen on exhibition in the Mineralogical Museum at Harvard University is estimated to weigh about sixty pounds of which perhaps twenty is due to fayalite. It is shown in the sketch, Fig. 1.

The most complete crystal broke away from the matrix but may be fitted back into its place and then appears to be about one-quarter of the whole individual. It measures five and one-half inches in the direction of the *b* axis, six inches parallel to *c* and one inch and a half parallel to the axis *a*. Its broken edges are cleavages; the faces are in part quite plane but the pinacoid curves into a vaguely developed prism form; the faces could be accurately measured with the contact goniometer yielding definitely the form shown in the drawing, Fig. 2. A second crystal, less complete, was thicker and probably larger in all its dimensions. A fragment of a third crystal separates the two larger individuals. Embedded in the

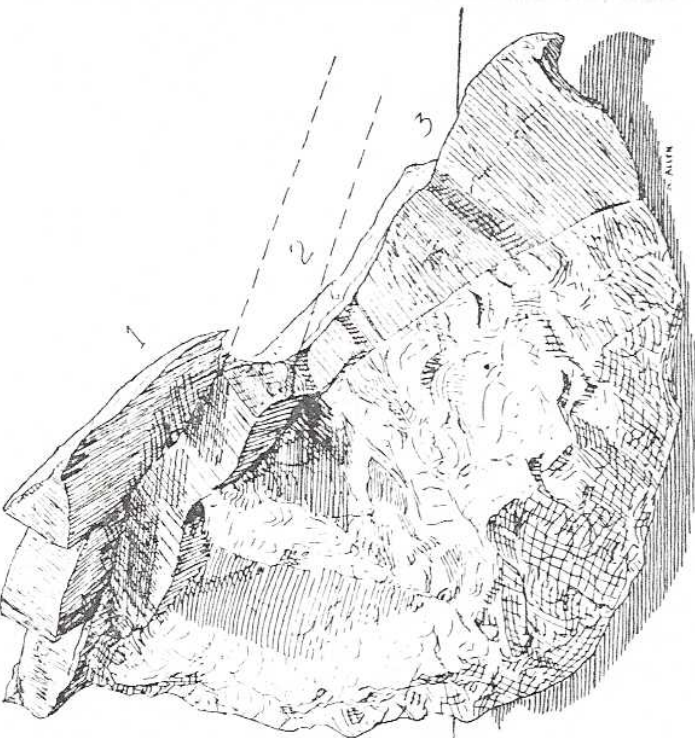


FIG. 1. Fayalite crystals in microcline. The three crystals are indicated. The termination of the uppermost complete crystal is not visible.

surface of the fayalite were a few much flattened grains of a black amphibole, not fibrous and of high index, probably hastingsite.

The latest description of fayalite in the Rockport area was by Warren and McKinstry (1924). They described the pegmatites and their minerals, and in several places refer to the last-described mass of pegmatite

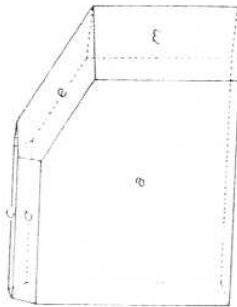


FIG. 2. Crystal of fayalite, $c\{001\}$, $a\{100\}$, $m\{110\}$, $d\{101\}$, $e\{111\}$. The prism is not plane on the crystal but curves into the pinacoid.

which was placed at their disposition. They were convinced, as I had been, that this mass was a xenolith in the Rockport granite. They also described a fourth occurrence of fayalite found in place by Dr. Warren on Brier Neck near Gloucester, on the opposite side of the Cape from the Babson Farm Quarry. This was in a dike-like mass of pegmatite which however is not well exposed and can not be proved to be an inclusion although they believe it to be one. This fayalite is in large part altered by a reaction with feldspathic material to a mixture of biotite, grunerite and magnetite.

The fayalite from all four of these occurrences has the same optical and physical properties. Each was found in an isolated mass of considerable size and in the case of the Babson Farm Quarry the evidence for the xenolithic nature of the mass was clear. It is perhaps safe to conclude that all were of the same nature. If this be true then the formation of fayalite on Cape Ann belongs to a period of granitic intrusion and crystallization preceding that of the present dominant regional rock, the Rockport granite. Such a rock is described by Warren and McKinstry under the name of nordmarkitic granite. This granite, which is cut by the dominant granite, contains fayalite as one of its lesser constituents, which is in harmony with the preceding conclusion as to fayalite pegmatite. No fayalite has ever, so far as the record shows, been found in the very numerous pegmatites characteristic of the latest Cape Ann granite.

With the decay of the quarrying industry on the Cape, most of the quarries have been abandoned and it does not seem likely that any further finds of these scattered blocks containing fayalite will be discovered.

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