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LUNAR SAMPLE STUDIES

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Prepared by NASA LYNDON B. JOHNSON SPACE CENTER



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Preface

This publication reports the results of five separate investigations of lunar samples which, while heralding no important new discoveries and theories in planetary science, contain nevertheless a wealth of descriptions and compilations valuable by themselves and important enough to be published by NASA. The samples and the nature of their analyses are:

An Apollo 15 breccia (15015), which is thoroughly analyzed as to the nature of the mature regolith from which it derived and the time and nature of the lithification process.

Two Apollo 11 and one Apollo 12 basalts (10069, 10071, and 12008), analyzed in terms of chemistry, Cross-Iddings-Pirsson-Washington norms, mineralogy, and petrography.

Eight Apollo 17 mare basalts (70017, 71055, 74255, 75075, 70215, 71569, 74275, and 75035), also analyzed in terms of chemistry, Cross-Iddings-Pirsson-Washington norms, mineralogy, and petrography. The first seven are shown to be chemically similar although of two main textural groups: the eighth is seen to be distinct in both chemistry and mineralogy.

A troctolitic clast from a Fra Mauro breccia (14321), analyzed and contrasted with other high-temperature lunar mineral assemblages. Two basaltic clasts from the same breccia are shown to have affinities with rock 14053.

And finally, the uranium-thorium-lead systematics of three Apollo 16 samples (60018, 60025, 64435) are determined; serious terrestrial-lead contamination of the first two samples is attributed to bandsaw cutting in the lunar curatorial facility.

The text, tables, and illustrations have been reviewed and approved for publication by a Lunar Samples Studies Editorial Board, chaired by William C. Phinney and Henry E. Clements.

February 1977

WILLIAM C. PHINNEY Lyndon B. Johnson Space Center Ť

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Contents

Chapter		Page
1	THE HISTORY OF LUNAR BRECCLA 15015 The European Consortium	1
2	COMPOSITION OF LUNAR BASALTS 10069, 10071, AND 12008 E. Jarosewich and Brian Mason	35
3	COMPOSITION OF EIGHT APOLLO 17 BASALTS Brian Mason, E. Jarosewich, Sara Jacobson, and G. Thompson	41
4	TROCTOLITIC AND BASALTIC CLASTS FROM A FRA MAURO BRECCIA N. G. Ware and D. H_Green_	49
5	U-Th-Pb SYSTEMATICS OF APOLLO 16 SAMPLES 60018, 60025, and 64435; AND THE CONTINUING PROBLEM OF TERRESTRIAL Pb CONTAMINATION OF LUNAR SAMPLES P. D. Nunes, D. M. Unruh, and M. Tatsumoto	61

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1. The History of Lunar Breccia 15015

The European Consortium and Friends^a

Breccia 15015, collected at the Apollo 15 lunar module station, is a coherent, tough soil breccia with an extensive glass surface coating. The wide range of data obtained on this rock is presented. The information concerns the nature of the mature regolith from which the rock was formed and the time and nature of the lithification process.

Lunar breccia 15015 is a coherent, tough breccia; 95 percent of the observable surface is glass coated. This large rock (fig. 1-1) (25 by 15 by 13 cm; 4770 g) was collected about 20 m west of the Apollo 15 lunar module (LM) site. It was first noticed by astronaut Scott during the standup extravehicular activity as unique in appearance and apparently unrelated to any nearby rocks or craters. Although 15015 was not documented during collection, its orientation on the lunar surface is known. A large portion of the rock, particularly the top and sides, is highly vesicular and cindery in texture.

The polarization properties of the rock indicate a ubiquitous vitreous coating on the exterior surfaces. The bulk chemistry of this glass is similar to the bulk composition of the rock and appear. on petrological grounds also, to have been derived from fusion of the rock. The rock contains a wide range of lithic fragments, among them both mare and potassium, rare-



Figure 1-1.- Highly vesicular glassy breccia 15015.

earth elements and phosphorus (KREEP¹) basalts, mineral fragments, and glasses, including devitrified "green glass." These components are typical of the Apollo 15 regolith in general and in detail are characteristic of the surface material at the LM station. Argon-40/argon-39 (40 Ar/ 39 Ar) ages indicate at least one clast older than 3.7 billion years (b.y.). The

^aThe team composition is listed in Acknowledgments at the end of this section.

¹Term used to express chemical composition of lunar materials, after llubbard, N. J.; and Gast, P. W.: Chemical Composition and Origin of Nonmare Lunar Basalts. Proceedings of the Second Lunar Science Conference, vol. 2, MIT Press (Cambridge, Mass.), 1971, p. 999.

LUNAR SAMPLE STUDIES

 38 Ar/ 37 Ar ages, noble gases, and track studies indicate an extensive preconsolidation exposure to cosmic rays, and an unusually large exposure age for one clast limits the consolidation age to less than 2.7 b.y. The "isochron age" of frothy vesicular glass tentatively indicates consolidation 1.0 b.y. ago. Carbon (C) data and noble gases show that the constituent grains have been heavily irradiated by solar wind. At some time in its history, possibly during impact lithification, 15015 experienced a nonuniform heating that caused light or more volatile species to be preferentrally lost from its upper surfaces. After formation, the precia was buried at a depth greater than 2 m until it was ejected onto the surface, possibly as recently as 30 million years (m.y.) ago.

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SAMPLE PROCESSING

The Consortium allocation, 15015,15 (150 g) was cut from the whole rock as a column to provide material from the top, inverior, and bottom of the sample (fig. 1-2). Aliquots of material from each of these locations were studied for their C chemistry, rare-gas content, cosmic ray fission tracks, traceelement composition, and selenochronology. In addition, surface samples from both top and bottom were investigated for their optical properties. Polished thin sections (15015,14,132 to 135) for the mineralogical and petrological studies were prepared from a slice adjacent to the surface sampled for the other measurements.

All sawing operations were performed at the University of Bristol with a water-cooled saw operating in a stainless steel nitrogen-filled glove box situated in a clean-air facility (ref. 1-1). The rock was fed slowly into a 15.2- by 0.05-cm diamond cutting disk by a counterbalance. Chipping operations were conducted directly in the flow of a laminar-flow clean-air bench using a stainless steel hammer and chisels. The nitrogen glove box and all materials coming into contact with the samples (aluminum foil, forceps, chisels, etc.) were precleaned by ultrasonic extraction in redistilled analytical reagent grade toluene methanol (3:1). All processes were documented photographically. The sample allocation plan is shown in table 1-I. The bulk chemical analysis (table 1-II. analysis 1) was performed on the dust generated during sample division.



Figure 1-2.- Sample subdivision of lunar breccia 15015.

The procedures used for the determination of methane (CH₄) and carbide (ref. 1-2), rare gases (ref. 1-3), bulk, trace chemistry, and electron microprobe analysis (ref. 1-4), track studies (ref. 1-5), seleno-chronology (ref. 1-6), and optical properties (ref. 1-7) have been detailed previously elsewhere. Total C, sulfur (S), and nitrogen (N) analyses were performed by submitting samples to complete combustion in a partial atmosphere of oxygen (O). Carbon was collected as carbon dioxide (CO₂). S as sulfur dioxide (SO₂), and N as nitrogen gas (N₂). During this procedure, an attempt was made to measure the volume of helium (He) released: however, the amount released was below the detection limit of the manometer used (0.002 ml, or 5 × 10⁻³ cc/g of analyzed sample).

Position of sample	Sample no.	Investigation
Top surface	15015,22	Tracke
	15015,23	Selenochronology
	15015,24	Optical properties
Surface	15015,26	Selenochronology
Clast	15015,27	Rare gaees
Тор	15015.16	Trace elements
	15015,17	Carbon, nitrogen, sulfur
	15015.19	Selenochronology
	15015,20	Rare Cases
	15015,21	Methane and carbide
Middle	15015,6	Methane and carbide
	15015,7	Carbon, nitrogen, sulfur
	15015,11	Trace elements
	15015,13	Selenochronology
	15015,14	Rare gas
	15015,29	X-ray photoelectron spectroscopy
Bottom surface	15015.8	X-ray photoelectron
	-	spect rosc opy
	15015,9	Optical properties
	15015,10	Tracks
	15015,28	Mineralogy/petrology
Bottom	15015,2A	Methane and carbide
	15015,2B	Trace elements
	15015,3	Carbon, nitrogen, sulfur
	15015,4	Rare gases
	15015,5	Selenochronology
Polished thin	15014.	Mineralogy/petrology
section	132 to 135	
	1	

TABLE 1-I.- SAMPLE ALLOCATION PLAN

Samples examined by X-ray photoelectron spectroscopy (XPS) were mounted on the sample holder of an ES200 spectrometer with epoxy adhesive in such a way that four faces could be independently examined. To avoid spurious contributions to the spectra, the adhesive region was covered with gold foil.

RESULTS AND DISCUSSION

Optical Properties

The top surface of breccia 15015 (15015.15.24) is a very-dark hued sample with a highly glassy. specularly reflecting surface. The geometric albedo (measured at a phase angle of 5°) is 0.070 at wavelength $\lambda = 350$ nm (3500 Å), 0.073 at 450 nm (4500 Å), and 0.083 at 600 nm (6000 Å). The amount of polarization ($P = (I_p - I_p)/(I_p + I_p)$) reaches a maximum of 0.415 at a phase angle of 120°, corresponding almost to the Brewster angle of silicates. These polarization properties are characteristic of speculat reflection from dark vitreous surfaces of complex structure.

The rate of change of polarization with phase angle (the so-called polarization slope), generally determined at phase angles near 30° , is related to the geometric albedo. In this respect, the extensive glassy top surface of 15015, 15, 24 conforms to the polarization slope/albedo law previously established for pulverized terrestrial samples, lunar fines, and rough lunar breccia surfaces. This law has been applied to the determination of asteroid diameters (refs. 1-8 and 1-9).

The exposed top surface of breccia 15015,15,24 is outstanding for the ubiquity of its black glass coating, generally smooth and vesicular, and possibly splashed on. Some parts of the thin shell exhibit cavities several millimeters in diameter. Freshly chipped surfaces are rough and do not produce glassy material.

Dimple microcraters, apparently not of impact origin, are abundant at the surface of the glass coating, as seen in figure 1-3. Some areas display a network of mounds (fig. 1-4) of almost equal size – a few tenths of a micron in diameter – and often regularly spaced or alined in chains. These mounds suggest metallic features similar to those postulated (ref. 1-10) to have been produced by reduction.

Chemical Analysis

Chemical analysis for major elements was performed on a homogenized sample, 15015,15, of powder and rock fragments obtained during sawing operations. A small amount of aluminum foil was the only visible contaminant, and this was removed by handpicking during the sieving of the sample.

Classical methods of analysis were used, as described by Scoon in reference 1-4. The values obtained are given as analysis 1, table 1-II, and they agree closely with the matrix samples analyzed by Rhodes (written communication) given as analyses 14 and 15, table 1-II. These values are in close agreement with the <1-mm fraction of the contingency soil sample 15021 from the LM site, analysis 13, table 1-II. This similarity in composition extends to the trace-element abundance determined by reference 1-11.

The composition is intermediate between mare basalts and the Fra Mauro basaltic glasses described by reference 1-12. These relationships are expressed

LUNAR SAMPLE STUDIES

TABLE 1-II. - CHÉMICAL ANALYSES AND CIPW NORMS OF BRECCIA 15015ª

[Analyses of whole rock and representative glass fragments]

(a) Description of samples

Analysis no.	Sample description
1	15015,15 whole rock analysis; analyst, J. H. Scoon
2	Glass; light brown, from vesicular external skin of 15015,133
3	Glass; light brown, smooth external skin of 15015,15 to 25
4	Glass; light brown, from vesicular vein penetrating sample
5	Glass; very pale brown, finely devitrified broken sphere; green glass spheres (GGS) ^b
6	Glass; somewhat inhomogeneous pale-brown striplike fragment molded on matrix; possibly a ''splash'' glass (mare basalts (M)) ^b
7	Glass; very pale green sphere (M)
8	Glass; brownish fragment with single vesicle (high-K Fra Mauro basalt (HK)) $^{ m b}$
9	Glass; brown-tinted, investing hypersthene crystal; a cored ''microwomb'' (HK) ^b
10	Glass; colorless or very pale green fragment of 3phere partially devitrified at edge; medium-K Fra Mauro basalt (MK) ^b
11	Glass; colorless sphere
12	Glass; colorless sphere; highland basalt (HB) ^b
13	15021 <1-mm fines; LM site; Apollo 15 Preliminary Examination Team .
14	15015,15,25 matrix samplé; analyst, J. M. Rhodes
15	15015,15,11, matrix sample; analyst, J. M. Rhodes

^aAnalysis 1 by classical wet procedure; analyses 2 to 12 by electron microprobe; analyses 13 to 15 by X-ray fluorescence.

^bClass types from reference 1-12.

graphically for certain major elements in figures 1-5 to 1-7.

Breccia 15015 may therefore be a shock-lithified representative of a regolith similar to 15021. Such a regolith could be predominantly derived from the local rocks of the Apollo 15 site, or, if the collection area lies on an ejecta ray, it could reflect in part compositions of rocks from a distant source area. In the former case, one would expect the block 15015 to have been ejected from some local crater whereas, in the latter case, the possibility exists that it was shock-lithified at some distant source and represents a residual boulder in fine-grained ray material deposited at the Apollo 15 site.

Mineralogy and Petrology

Rock 15015 is probably a shock-fused polymict soil breccia. The matrix is composed of glass or its cryptocrystalline devitrification products crowded with minute (1 to 2 μ m) mineral fragments in which pyroxene predominates over plagioclase. The glass is gray-brown in sections <15 μ m thick but almost black and opaque in thicker sections. The opacity is in part due to submicroscopic iron (Fe) or troilite droplets and "ilmenite" crystallites dispersed in the glass.

Larger rock, mineral, and glass fragments are set in this base. Most of the fragments fall within the fol-

TABLE 1-II. - Continued

(b) Chemical analysis

Compound		Analysis no. ^d , wt.Z													
	T	2	3	4	5	6	7	8	9	10	11	12	13	14	15
510,	47.11	48.01	48.49	47.45	45.65	47.46	47.03	55.30	50.29	49.74	48.61	46.66	40.55	47.50	47.30
TIOZ	1.90	۲.89	1.80	1.92	. 37	2.02	2.20	2.16	1.94	1.56	.58	. 38	1.75	1.76	1.74
A12 ⁰ 3	14.46	13.43	13.96	13.62	7.53	10.31	12.15	14.16	16.12	16.64	17.61	25.91	13.73	13.99	14.16
Cr203	.40	.02	.40	. (e)	.60	.51	.41	.12	.15	.25	. 32	.22	(e)	(e)	(e)
FeO	14.18	12.76	14.37	14.94	20.44	19.50	18.08	11.10	10.66	9.49	8.46	5.28	15.21	14.63	14.56
MgO	.19	(e) 10.16	9.56	9.50	.31	.32	8.27	.17	.16 6 39.	.17	.16	.08	.20	.21	.22
Cal	10.47	11.16	10.41	9.94	8.25	10.36	11.06	8.91	10.25	10.56	10.82	10.24	10.54	10.52	10.62
Na20	.31	. jó	.51	.51	.23	.30	.25	1.06	. 82	.76	.06	.17	.41	.50	.50
s20	.28	.24	.29	.01	.02	.13	.05	1.24	.95	.49	.02	.00	.20	.24	.22
^P 2 ^O 5	.22	.05	(e)	(e)	(e)	(e)	(e)	. 33	.47	. 36	(e)	(e)	.18	.24	.22
Total	99.65	98.28	100.02	97.91	101.14	102.43	99.77	99.25	98.69	99.26	100.95	96.42	99-15	99.71	99.65

TABLE 1-II. - Concluded

(c) CIPW norm

Mineral	Analysis no. ^d , wt.2									_					
	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15
Quartz Orthoclase Albite Anorthite (An) Plagioclase Diopside Hyperstheme Olivine Chromite Ilmenite Apaite Molt Feo+mgD	 1.65 4.32 36.34 40.65 11.69 38.22 2.86 .59 3.61 .47 4, 60	0.35 1.39 4.74 38.18 17.77 36.86 	0.95 1.34 4.20 35.11 39.31 13.69 40.52 .60 3.41 42.9	1.04 .06 4.32 34.84 39.16 11.98 42.04 3.65 	0.12 1.95 19.45 21.40 17.66 27.13 33.24 .88 .70 	0.77 2.54 26.40 28.94 20.77 37.15 10.22 .75 3.84 	0.83 .30 2.12 31.88 34.00 19.26 40.57 .60 4.18 52.40	14.02 7.33 8.97 30.22 39.19 9.91 23.76 .18 4.10 .78 52.50	7.13 5.61 6.94 37,50 44.44 5.81 29.55 	3.27 2.90 6.43 40.55 46.98 7.74 34.23 .37 2.96 .85 33.00	 0.12 .15 47.72 48.23 4.80 41.85 4.38 .47 1.10 23.80	 1.44 69.93 71.37 4.59 21.74 2.68 .32 .72 26.90	1.18 3.47 35.03 38.50 13.30 36.31 6.05 3.32 .43 42.70	1.47 4.23 35.22 39.45 12.75 39.44 2.69 	1.30 4.23 35.74 39.97 12.81 38.15 3.30 .32 42.20
Noll An ir norm plagioclase	88.80	86.90	8 0	88.40	90 .90	90.70	93.80	76.00	83.60	85.60	98.90	97.90	90.49	88. 70	88.64

 $^{\rm A}$ Analysis 1 by classical wet procedure; analyses 2 to 12 by electron microprobe; analyses 13 to 15 by X-ray fluorescence.

^bGlass types from reference 1-12.

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d From table 1-11(a).

Not determined.

fCross-Iddings-Pirsson-Washington norma.

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lowing size ranges: minerals, 0.02 to 0.3 mm; glass fragments, 0.02 to 1.0 mm; lithic fragments, 0.20 to 1.50 mm. Within the lithic fragments, basalts of several types predominate (84 percent) over metaclastic rocks (16 percent). Sporadic lithic fragments of basaltic composition up to 1 cm in size also occur. The distribution of fragment types is somewhat variable as is seen from the modes of four thin sections given in table 1-III. No distinct banding or segregation can be observed.

One surface of the rock, seen in section 15015,133, is coated by a vesicular glass of cindery appearance. The surface of the rock appears frothy and partially melted to a depth of about 5 mm. In thin section, every surface of the rock is seen to be coated with a thin skin of light-brown-colored glass that, with an increasing amount of resorbed mineral fragments, merges in an irregular manner into the

body of the rock. On the opposite surface, the glass 15015,15 to 25 is more in the nature of a glazed skin up to 0.5 mm thick coating the rock. In thin section, the glass is similar in color but less vesicular than the frothy glass. Flow bands are visible in thin section and are accentuated by minute submicron droplets of metallic Fe and troilite. The contact between the glass skin and the rock is smooth, and the transition from one to the other is clear-cut.

The composition of the two glasses is quoted in table 1-11, analyses 2 and 3, along with the composition of a thin vein penetrating the sample (analysis 4). These compositions are similar and close enough to the composition of the whole rock to suggest that they have developed from the melting of the whole rock or of a slightly heterogeneous soil or rock of closely allied composition (figs. 1-5 to 1-7).



Figure 1-3.- Dimple craters on breccia 15015,15,24.

MERODUCIBILITY OF THE MALE PAGE IS POOR



Figure 1-4.- Micromounds on breccia 15015.15,24.

Component	Volume, percent								
	Polish	ed thin	section	no.	Average				
	132	133	135	142					
Rock fragments Pyroxene fragments Plagioclase fragments Glass fragments Metal, oxide, sulfide fragments Matrix	16.1 16.3 5.6 9.4 1.1 51.1	8.8 12.2 7.2 12.7 2.2 56.9	8.6 26.0 8.2 9.1 2.4 45.7	8.0 21.9 10.4 8.9 .4 50.4	10.4 19.1 7.8 10.0 1.5 51.2				

TABLE 1-III	MODAL	MINERALOGY	OF	BRECCIA	15015
-------------	-------	------------	----	---------	-------



- + Averages of Apoilo 15 glass lypes (ref. 1-12)
- Three analyses of rock 15015
- Bubbly coating glass on 15015
- ♥ Glass seam in 15015, probably splash glass
- Individual glass fragments
- M < 1 mm fines, mare sites

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- F >1 mm fines, Apennine front sites
- Figure 1-5.- 15015 glass fragments and coatings, ferrous oxide/magnesium oxide (Fe0/Mg0). GG = green glass, HB = highland basalt, HK = high-K Fra Mauro basalt, LK = low-K Fra Mauro basalt, MK = medium-K Fra Mauro basalt, and M1 to 4 = mare basalts 1 to 4.

The principal difference between the veining and coating glass and the whole rock is that the latter is just olivine normative (2.5 to 3.5 percent) and the former just quartz normative. Such differences, if the possibility of analytical error is excluded, could be reasonably associated with a minor heterogeneity in the original rocks or with incomplete melting in the production of the coating glass. Molten material might possibly have been added in flight in an impact-produced cloud that was subsequently laid down as a ray deposit. The latter view would be more in accord with the studies of Carter (ref. 1-13), who made a detailed scanning electron microscope (SEM)



- Glass seam in 15015, probably splash glass in original deposit
- Individual glass fragments
- Figure 1-6.- 15015 glass fragments and coatings, ferrous oxide/alumina (Fe0/Al₂0₃).



- + Averages of Apollo 15 glasses (ref. 1-12)
- Three analyses of rock 15015
- Bubbly coating glass on 15015
- Glass seam in 15015, probably colash glass
- Na₂O+K₂O in individual glass fregments
- P₂O₅ in individual glass frágments
- Figure 1-7. 15015 glass fragments and coatings. Composite plot of sodium oxide plus polassium oxide (Na $_20 \div K_20$) and phosphorus pentoxide (P $_20_5$) against alumina (Al $_20_3$). The lines join analyses of individual glass fragments in which the three oxide contents have been determined.

study of the exterior glass from sample 15015.36. He described metallic Fe and troilite mounds, whisker structures, outgassing structures, and low-velocity, hot-target impact structures on the glass surface. The glass coating was interpreted as a siliceous melt generated in a hot, impact-produced cloud. In such a situation, strict comparability of the composition of coating glass and coated rock would not be expected. Reference 1-13 also indicated that the absence of high-velocity, microimpact craters on the glass examined suggests that this sample was from a surface that had been buried or shielded while on the Apollo 15 site.

Mineral Clasts

Mineral clasts are angular to subangular fragments the maximum grain size of which may be up to three times that found in the same minerals in the lithic clasts. This finding indicates that rocks coarser in grain size than the lithic clasts or the degradation products of such rocks existed in the source areas of the material that contributed to the makeup of rock 15015.

Most of the mineral clasts show evidence of shock metamorphism in their undulose extinction, irregular lamellar features, and, in the case of plagioclase, the presence of felsitic devitrified maskelynite. Rare fragments, usually hypersthene, are coated with a thin layer of brownish glass; this glass has affinities with the Fra Mauro basaltic glasses of reference 1-i2. One example is quoted as analysis 9, table 1-11.

Pyroxene clasts cover a wide range of compositions that embrace those of the pyroxenes of the lithic clasts. Analyses of typical examples are quoted in table 1-IV and plotted on the pyroxene quadrilateral in figure 1-8. It is possible that the hypersthenes with >85 percent enstatite (En) are derived from coarsely crystalline rocks of the anorthosite, norite, troctolite (ANT) suite that are proposed here as a source of the plagioclase clasts with approximately 96 percent anorthite (An).

Plagioclase clasts range in composition from 97 to 80 percent An with orthoclase (Or) content from 0.1 to 0.8 percent in the more sodic members. Normally, the clasts are not zoned; a histogram of compositions is given in figure 1-9, along with the composition of the plagioclase in the lithic clasts studied. A popula-



- Fra Mauro-type basalts
- Metamorphic (recrystallized) rocks, probably metaclastic, some Fra Mauro, others ANT suite
- + Single-crystal clasts



tion of plagioclase clasts is distinctly richer in An than the feldspars in the lithic clasts. This group with approximately 96 percent An is probably derived from more coarsely crystalling anorthosis crocks of the ANT suite that were not represented among the lithic clasts. The group with 88 to 93 percent An is probably derived from the mare basalts and the feldspathic basalts recognized among the lithic clasts; the group with approximately 88 percent An may have a source in the disintegration of the Fra Mauro basalts recognized among the lithic clasts.

The ferrous oxide (FeO) content of the plagioclase clasts shows some correlation with An content. The majority of the clasts in the range of 90 to 97 percent



Composition of plagioclase in lithic fragments



LÚNAR SAMPLE STUDIES

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Compound,	·			Analysi	B		
element, or mineral	1	2 3		4	5	6	7
	ťýp	ersthene		Subcalcic augite	Subcalcic ferroaugite	Augite	Ferroaugite
510 ₂	54.70	53.76	51 .8 6	49.78	48.86	50.10	47.88
T10,	.18	1.31	.25	.92	.64	.80	' 1 _~ 26
A1,0,	4.54	1,94	.97	2.04	1.89	2.48	1.10
Cr ₂ 0 ₃	. 51.	. 33	.74	.58	.74	1.11	.22
řeO MaO MgO	8.80 .15 30.26	9.68 .90 30.42	14.88 .29 26.28	22.99 .42 14.40	29.12 .61 10.56	17.12 .28 11.65	28.15 .46 6.57
CaO	1.55	1.22	1.81	8.10	8.14	16.24	13.75
Na ₂ 0	.00	.00		* زانلہ . مرد	.02		•••
к ₂ 0	.00	.01		.00	.01	••	
Total	100.71	99.57	97.08	99.23	100.59	99.78	99.39
Enstatite (En) Ferrosilite (Fs), mol2	85.7 13.9	81.6 16.0	73.2 23.2	43.2 39.4	31.9 50.4	35.3 29.2	20.2 4°.4
Wollastonite (Wo)	2.3	2.4	3.6	17.4	17.7	35.5	30.4
			Unit for	mulas			
Silicon (Si) Titanium (Ti)	1.907	1.911 .036	1,939 .008	1.929 .027	1.926 .019	1.922	1.937 .039
Aluminum IV (A1 ⁻)	.093	.082	.043	.071	.074	.088	.053
Aluminum VI (Al ^{VI}) Chromium (Cr)	.094 .014	.010	 .022	.021 .018	.014 .024	.024 .034	.08
Irón (Fe) Managanaga (Ma)	.257	.288	.466	.745	.960	.550	.953
Magnésium (Mg)	1.573	1.612	1.465	.832	.021	.010	. 397
Calcium (Ca)	.044	.047	.073	. 337	.344	.668	.596
Sodium (Na)	.000	.000		.000	.002		
rotessium (K)	.000	.001	••	.000	.001		••

TABLE 1-IV. - ELECTRON MICROPROBE ANALYSES OF BRECCIA 15015 FYROXENE CLASTS

An contain less than 0.1 percent FeO, whereas clasts in the range of 80 to 90 percent An show higher concentrations of up to 0.9 percent FcO.

T TINE

TAL SUCTION

Metallic Fe occurs as sporadic rounded fragments up to 50 μ m in diameter. The fragments fall into two subgroups: (1) fragments with <2.0 percent nickel (Ni), 0.1 to 0.6 percent cobalt (Co) and <0.1 percent phosphorous (P), probably derived from basaltic rocks and (2) fragments with 3.0 to 10.0 percent Ni, 0.3 to 0.5 percent Co, and <0.2 percent P, probably of meteoritic origin.

Vitric Clasts

Vitric clasts constitute approximately 10 percent of rock 15015. They include indeterminate plastic forms, twisted and ropey forms, spheres, and broken fragments. One unique clast had the appearance of a splash glass in thin section and a composition similar to that of a mare-type basalt (analysis 6, table 1-II).

The glass fragments may be completely fresh or show partial, usually marginal, devitrification. One type, chemically similar to the green glass spheres described by reference 1-14, is always completely devitrified.

Five main chemical categories of glass fragments may be recognized and can be correlated with the glass types recognized by reference 1-12. Representative analyses of these types are quoted in table 1-II, and these and other glasses from 15015 are plotted in figures 1-5 to 1-7 along with the average compositions of the glass types recognized in the Apollo 15 soils by Reid et al. (ref. 1-12).

The similarity between the bulk composition of 15015 and contingency fines 15021 has already been mentioned. Although fewer glass analyses are available from 15015, it will be seen from comparison of figures 1-5 and 1-6 with figures 2(a) and 2(b) of reference 1-12 that the general similarity extends to the compositions and relative abundances of the glass fragments in the two samples.

These five main chemical categories of glass fragments are grouped as follows.

1. Fra Mauro basaltic glasses are the most abundant: they consist of ropey, plastic, and broken forms; in thin section ranging in color from very pale fawn to pale brown. No glass spheres were observed. Low-, intermediate-, and high-K types are represented. Compared with the contingency fines, 15015 contains a higher proportion of high-K members that are slightly richer in Fe than those of 15021. Phosphorus pentoxide (P_2O_5) has not been determined in all the glass analyses, but in figure 1-7 it is seen that glasses with high sodium oxide plus potassium oxide ($Na_2O + K_2O$) often have high values of P_2O_5 , which is a characteristic of Fra Mauro, or KREEP glasses.

2. Agglutinates are a minor group; they tend to be heterogeneous and to have approximately the same composition as the bulk of 15015.

3. Mare-type basaltic glasses are a subordinate group. They usually occur as fragments and occasional spheres with a yellow-brown to green-brown color. Too few have been analyzed to allow the recognition of any significant subdivisions within them.

4. Highland basalt glasses – a few colorless fragments and spheres have been analyzed with the high alumina (Al_2O_3) and calcium oxide (CaO) and low FeO characteristic of this group.

5. Green glass spheres occur as a subordinate glass type but are always represented by devitrified spheres or broken forms composed of fawn-colored cryptocrystalline devitrification products. They form a tight cluster close to the average determined by reference 1-12 for the green glass spheres. This may be seen in figures 1-5 to 1-7.

Such devitrified types of the unique green glass are not uncommon in soils and breccias of the Apollo 15 site. Reid et al. (ref. 1-12) comment on their abundance along with unaltered spheres in 15601; in reference 1-15, their presence is noted as a minor component in 15425 as a green clod dominantly composed of the green glass.

The gteen glass spheres conceivably are the products of several events operating on identical material. However, their uniformity in composition suggests their formation in a single event, irrespective of magmatic, meteoritic, or impact origin.

The presence of devitrified green glasses implies that annealing occurred in some thermal event subsequent to their initial formation. The event associated with this annealing could have been a secondary

cratering (if 15015 were shock-lithified near the Apollo 15 si.e) or a primary event if shock-lithification were associated with ejection in the Aristillus or Autolycus tay, of which 15015 may be an integral component. If the green glass is a component of the ray, then it must also either have been a component of the regolith of the Aristillus-Autolycus area or have been generated in the major-impact explosion and devitrified by a secondary cratering event acting on the deposited ray material.

Lithic Clasts

Lithic clasts constitute approximately 10 percent of rock 15015. The following types are recognized: mare basalts (30 percent), F1a. Mauro basalts (33 percent), feldspathic basalts (22 percent), and metamorphic (recrystallized) clasts (15 percent).

Mare basalts.— Mare basalts are represented by fragments of porphyritic clinopyroxene-tich basalts, potphyritic pyroxene vitrophyres, and subophitic ilmenite basalts. Insofar as 1-mm fragments are valid for an assessment, these basaltic fragments can be matched texturally with mare basalts collected at the Apollo 15 site. It is significant that olivine-bearing types are rare among the fragments in 15015.

Analyses of minerals from these typical fragments are quoted in table 1-V(a), analyses 1 to 8, and the pyroxene compositions are plotted on figure 1-8. The pyroxenes are commonly zoned and range from early formed, pale-colored, calcium-poor pigeonite with a ratio of Fe to Fe plus magnesium (Mg) approximately 0.30 by way of calcium-rich pigeonites to late ferroaugite with Fe/Fe+Mg approximately 0.55. The late pink-brown ferroaugite may contain up to 2.0 percent titanium dioxide (TiO₂) and 9 percent Al₂O₃.

Fra Mauro basalts. – Fra Mauro basalts are represented by mesostasis-rich (or intersertal), olivine-free basalts with up to 45 percent lathlike, often slightly curved, crystals of plagioclase. The plagioclase in the fragments analyzed is restricted to compositions of approximately $An_{83.5}$, albite (Ab) 15.0, $Or_{1.5}$.

The pyroxenes are zoned from colorless to very pale yellow-brown hypersthene with Fe/Fe+Mg approximately 0.15 to pale yellow-brown calcium-poor plgeonites with Fe/Fe+Mg approximately 0.35. Occasional Fe-rich ferroaugite was observed. Early formed hypersthenes are relatively aluminous compared with the later crystallizing hypersthene and pigeonite. Where a fine-grained mesostasis is present, it is composed of skeletal crystals of ilmenite, pyroxene, and plagioclase with interstitial potash feldspar, cristobalite, and troilite. No quantitative data were obtained from this material.

Analyses of minerals from four typical clasts are quoted in table 1-V(b), analyses 25 to 38, and the pyroxene relationships are plotted in figure 1.8.

Feldspathic basalts. - The feldspathic basalts include fragments that were impossible to assign to either of the two principal groups just described. They are more feldspathic than the unambiguous pyroxene-rich mare basalts, but their plagioclase has a higher An content and their pyroxenes a higher Ee/ Fe+Mg ratio than the basalts here classified as of Fra Mauro type. The feldspathic basalts include variolitic, intersertal, and subophitic types of variable grain size in which olivine-bearing types are rare. A common type of clast consists of three or four subparallel crystals of tabular plagioclase with a few interstitial crystals of allotriomorphic pyroxene and minor ilmenite. The plagioclase constitutes 30 to 55 percent of the fragments, and its composition lies within the following ranges: An₈₆ to An₉₂, Ab₁₃ to Ab₇, Or_{0.2} to Or_{0.8}. In some clasts, plagioclase shows normal zoning; in others, it is unzoned. Pyroxenes constitute from 45 to 70 percent of individual clasts, and reference to table 1-V(a), analyses 9 to 24, and to figure 1-8 illustrates their wide range of composition. They range from calcic pigeonites with Fe/Fe+Mg approximately 0.35 to subcalcic ferroaugites or ferroaugites with Fe/Fe+Mg approximately 0.85. Essentially, the pyroxenes cover the field shown by the pyroxenes in the mare basalts of Apollo 15. However, the interpretation of trends in individual clasts is ambiguous. Analyses 10 to 12 (table 1-V(a)) are not unlike those of late-crystallizing pyroxenes of 15085,14, a mare basalt described in reference 1-16, but their trend is also similar to that of the late-crystallizing portion of 14310, which is a Fra Mauro basalt (ref. 1-17).

Gleadow et al. (ref. 1-18) applied chemical discriminant analysis to recognized petrographic types of clasts in Apollo 14 and 15 breccias, including 15015,19. They recognized a distinct field for mare and Fra Mauro basalts but not for a feldspathic group. Most of the feldspathic basalts here recognized

THE HISTORY OF LUNAR BRECCIA 15015

probably have closer affinities with mare basalts and represent olivine-free types with early crystallizing plagioclase.

Metamorphic (recrystallized) clasts. – Two different types of metamorphic (recrystallized) clasts are recognized on the basis of texture.

1. Feldspathic microgranulites are composed of polygonal crystals of plagioclase associated with subordinate granules of pyroxene with or without minor olivine and ilmenite. These may range from almost pure anorthosite to types with only 50 percent plagoclase. They appear to have been produced by thermal recrystallization of an already crystalline rock with no marked differences in grain size.

2. Porphyroclastic rocks have angular porphyroclasts of plagioclase approximately 100 μ m in size set in a finer grained matrix of plagioclase and hypersthene with or without minor olivine and ilmenite. The matrix grain size ranges from 5 μ m in the least to 30 μ m in the most recrystallized examples. Where

TABLE 1-V .- ELECTRON MICROPROBE ANALYSIS OF MINERALS FROM LITHIC CLASTS OF BRECCIA 15015

(a) Description of Samples

nalysis no.	Minèral		Description
	Mare-type, pyroxene-	rich	basalts with <30 percent plagioclase ⁸
1	Pyroxene - pigeonite core	1	Porphyritic pyroxene-rich basalt (135I)
2	Pyroxene - augite rim	1	
3	Pyroxene - pigeonite core	١	Porphyritic pyroxene vitrophyre (135F)
4	Pyroxène - augite rim	Ì	Pyroxene, 46 percent; mesostasis, 54 percent
5	Plagioclase)	Pyroxene-rich ilmenite basalt (135K)
6	Pyroxene	1	Pyroxene, 83 percent; plagioclase, 17 percent
7	Pyroxěne	(
8	Ilmenite	1	

Basalts with >30 percent plagioclase^b

9 10 11 12 13	Plagioclase Pyroxeme - pigeonite core Pyroxene - augite intermediate Pyroxene - ferroaugite interstitial Ilmenite	Coarse intersertal basalt (135L) Pyroxene, 53 percent; plagioclase, 47 percent
14 15 16	Plagioclase Pyroxene – core Pyroxene – interstitial	Intersertal basalt (133F) Pyroxene, 54 percent; plagioclase, 46 percent
17 18 19	Plağioclase Pyroxene – core Pyroxene – rim	Pyroxene-rich basalt (133D) Pyroxene, 69 percent; plagioclase, 31 percent
20 21	Plagioclase Pyroxene	Fine-grained basalt (135AB) Pyroxene, 45 percent; plagioclase, 55 percent
22 23 24	Pyroxene • subcalcic ferroaugite Plagioclase Ilmenite	Dark variolitic basalt (15,5B) Pyroxene, 50 percent; plagioclase, 40 percent; Ilmenite, 3 percent; mesostasis, 7 percent

^aAn, 88 mo1%; FeO, 1.0 wt.%.

^bAn, 92 to 87 mol%; FeO, 0.6 to 1.4 wt.%.

LUNAR SAMPLE STUDIES

pyroxene is dominant, it occurs as idioblastic rods set in interstitial plagioclase crystals of irregular outline. In the more feldspathic types, plagioclase forms (dioblastic plates, ard the pyroxene forms the interstitial crystals.

These porphyroclastic rocks were evidently formed by the recrystallization of fine-grained clastic rocks — including soils. The recrystallization is due to the heat remaining after shock-vitrification (or melting) of originally unconsolidated material. Increase in grain size and changes in texture of the matrix reflect both the temperature attained and the duration of the annealing period. .

On the basis of plagioclase composition, the clasts can be tentatively assigned to one of two chemical groups. Those with plagioclase with approximately 93 percent An are possibly recrystallized rocks in which members of the ANT suite or its fragmental associates predominate; e.g., analyses 44 to 47, table 1-V(b). Those with plagioclase approximately 86 percent An

TABLE_1-V.- Continued

(a) Concluded

alysis no.	Mineral-		Description
	Нурега	thene	"Fra Mauro" basalts ^C
25 26 27 28	Plagioclase Pyroxene - córe Pyroxene - rím Pyroxene - interstitiál	{	Mesostasis-rich basalt (133A) Pyroxene, 51 percent; plagioclase, 32 percent; mesostasis, 17 percent
29 30 31	Plagioclase Pyroxene - hypersthene Pyroxene - pigeónite	}	Mešostāsis-poor basalt (135M) Pyroxene, 60 percent; plaģioclase, 31 percent; mesostasis, 9 percent
32 33	Plagioclase Pyroxene – hypersthene	}	Pyrcxene, 30 percent; plagioclase, 38 percent; mesostasis, 32 percent (135H)
34 35 36 .17 38	Pyroxene - hýpérsthene Pyroxene - pigeonite Plagioclase Crístobálite Ilmenite		Pale, merostasis-rich basalt (15,238) Pyrožene, 30 pércent; plagioclase, 20 percent; mesostasis, 45 percent; ilmenite, 5 percent

Metamorphic (recrystallized) rocks, probably metaclastic"

39	Olivine	}	Hypersthene-rich betaclastic (135J)
40	Pyroxene - hypersthene		Femic minerals, 74 percent; plagioclase,
41	Plagioclase		26 percent
42 43	Olivine Pyroxene • hypersthene	}	Olivine, hypersthene, plagioclase granulite (135B4) Femic minerals, 49 percent; plagioclase, 51 percent (An ₈₅ Ab ₁₄ Or ₁)
44 45	Plagioclase Pyroxene - salite	}	Anorthositic granulite (133E) Femic minerals, 19 percent; plagioclase, 81 percent
46	Plagioclase	}	Noritic metaclastic (135G)
47	Pyroxene - hypersthene		Pyroxene, 52 percent; plagioclase, 48 percent

^CAn, 82 to 84 mol%; FeO, 0.4 to 0.6 wt.%.

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^dNumbers 39 to 43: Fra Mauro affinity; An, 85 to 88 mol%; Fe0, 0.3 to 0.4 wt.%. Numbers 44 to 47: ANT affinity; An, 92 to 94 mol%; Fe0, 0.1 to 0.3 wt.%.

Compound					Ana	lysis no., ^{a.}	wt.Z					
or element	1	2	3	4	5	6	7	8	9	10	11	12
\$10 ₂	£3.08	49.39	50.93	48.56	48.26	50.85	51.31	(b)	45.28	50.45	50.08	48.10
T10,	.22	1.17	.72	2.01	.02	1.05	.80	51.68	.02	.66	.85	1.10
A1203	2.49	6.53	5.78	9.21	33.16	3.31	1.34	1.27	32.73	2.71	2.63	1.45
^{Cr} 2 ⁰ 3	(b)	(b)	(b)	(b)	(ь)	(b)	(b)	(b)	(b)	(ь)	(b)	(b)
FeO	17.95	16.25	16.87	15.93	1.08	15.77	18.50	46.03	,68	23.21	21.72	30.49
Mm0	(b)	(b)	(b)	(Ъ)	(b)	(b)	(b)	(b)	(Ġ)	(ь)	(Ъ)	(Ъ)
MgO	22.26	13.49	19.65	13.61	.21	15.15	18.85	.00	.07	15.93	12.91	5.11
CaO	3.46	14.64	5 . 13	11.68	17.86	13.72.	6.88	(b)	18.97	6.25	10.74	12.96
Na ₂ 0	.02	.06	.07	.08	1.28	.05	. 30	(b)	,89	.07	.09	.04
₹ ₂ 0	.02	.02	. 02	.02	.05	.02	.02	(b)	.04	.02	.02	.02
Tetal	99.50	101.55	99.17	101.10	101.92	99.92	98.00	98.98	98.68	99.30	99.04	99.27
					Ur	nit formulas						
51	1.957	1.835	1.885	1.792	2.184	1.911	1.958		2.127	1.933	1.938	1.957
T1	.007	.033	.021	.056	.001	.030	.023	0.985	. 001	. 020	. 02 5	.034
AI IV	.043	. 165	.115	.208	1.769	.089	.042	.038	1.813	. 067	. 062	.043
NI ^{VI}	.066	. 122	.138	. 193		.058	.019			. 056	.059	.027
Cr	••											
Fe	. 554	.505	.522	.492	.042	.496	.593	.975	.027	.774	.703	1.038
to I	••									T-	••	
1 8.	1.223	.747	1.084	.749	.015	. 849	1.073	.000	.005	910	.745	. 310
Ca	.137	.583	.204	.462	.866	.553	.282		.955	257	.446	. 566
Na	.002	.005	.005	.007	.133	.004	.003		. 081	006	.007	.004
K	.002	.0 02	.002	.002	.003	.002	.002		. 003	. 002	.002	. 002
					4	tomic ratios	; ;					
	Mg 63.9	Mg 40.7	Mg 59.9	Mg 44.00	Ca 88.2	Mg 44.7	Mg. 55.1	Mg 0.00	Ca 91.9	Mg 47.6	Mg. 39.3	Mg 16.2
	Fe 28.9	Fe 37.5	Fe 28.8	Fe 28.9	Na 11.5	Fe 26.1	Fe 30.4	Fe 100.00	Na 7.8	Fe 38.9	Fe 37.1	Fe 54.2
	Ca 7.2	Ca 31.8	Ca 11.3	Ca 27.1	K .3	Ca 29.2	Ca 14.5		K .3	Ca 13.5	Ca 23.6	Ca 29.6

TABLE 1-V.- Continued

(b) Chemical analyses, unit formulas., and atomic ratios

a From table 1-V(a).

^bNot determined.

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TABLE 1-V.- Continued

(b)	Continued
(0)	oomernace.

Compound	Î				Anal	lysis no., ^a u	/t.Z					
or element	13	14	15	16	17	18	19	20	21	22	23	24
s10,	(b)	45.99	48.81	50.14	46.56	50.97	46.23	47.75	48.82	51.53	47.99	(ь)
T10,	52.90	.08	1.05	1.01	.08	.46	1.25	j .00	1.12	.26	(b)	51.15
A1,0,	.13	34.02	2.56	1.32	31.93	1.16	1.16	33.17	2.42	.79	31.51	. 37
2 J Cr ₂ 0	(ь)	(Ъ)	. 78	.40	(ь)	.87	.02	(ь)	(ь)	.19	(ь)	. 12
23 Feù	46.17	.60	18.25	26.66	1.03	19.61	36.69	1.36	28.09	28.31	.99	46.27
(m)	(b)	(Ъ)	.40	.05	(b)	.35	.48	(b)	(b)	. 17	(b)	. 06
Ng.)	.11	. 38	14.28	13.15	.01	20.49	3.99	.15	10.08	7.85	. 36	.81
Cảo	(b)	17.89	13.88	6.86	17.75	5.38	10.32	17.09	7.88	9.46	16.63	(Ь)
Na ₂ 0	(ь)	.99	(Б)	(Ь)	1.15	(Ъ)	(Ъ)	1.23	.25	(b)	1.43	(b)
R20	(b)	.04	(b)	(b)	.12	(b)	(b)	.02	.03	(ь)	.08	(ь)
Total	99.31	99.99	100.01	99.59	98.63	99.29	100.14	100.77	98.69	98.56	98.99	98.78
					U	nit formulas			•			
Si		2.122	1.873	1.956	2.180	1.927	1.920	2.183	1.943	2.052	2,228	
Ti	.007	.003	.031	.030	.003	.014	.040		. 034	. 008		0.98
A1 IV	. 004	1.849	. 116	.044	1.762	. 052 [.]	.058	1.788	.057		1.725	. 0
A VI			024	017					057	017		
A1 (-			.024	.017		026	001			. 037		00
CI Fo	97/	026	586	870	041	620	1 274	052	935	.963	039	
10 10			.014	.002		.012	.017			.006		.0
Mg	.005	. 026	. 817	.764	.007	1.155	.027	.011	.598	. 466	.025	.0
Ca	••	.885	. 571	.287	.891	.218	.460	.837	. 337	. 404	.828	
Na		. 089			. 105			.110	.020		.129	
K	••	.003		••	.0 08	•-		. 002	. 002		. 605	
						Atomic ratio	5					
	Mg 0.5	Ca 90.6	Mg 41.4	Mg 39.8	Ca 88.7	Mg 58.0	Mg. 12.4	Ca 88.2	Mg 32.0	Mg 25.7	Ca 86.1	Mg 3.
	Fe 99.5	Na 9.1	Fe 29.7	Fe 45.3	Na 10.5	Fe 31.1	Fe 64.3	Na 11.6	Fe 50.0	Fe 52.0	Na 13.4	Fe 97.
		к.3	Ca 28.9	Ca 14.9	к.8	Ca 10.9	Ca 23.2	к.2	Ca 18.0	Ca 22.3	K .5	

^aFrome table 1-V(a). ^bNot determined.

MUCIBILITY OF THE

LUNAR SAMPLE STUDIES

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TABLE 1-V.- Continued

Compound or element					Analysis no., ^a wti Z											
or element	25	26	27	28	29	30	31	32	33	34	35	36				
sio ₂	47.97	53.80	52.84	47.98	48.62	52.44	52.83	48.31	53.43	55.19	53.13	49.36				
Ti0,	.10	.57	. 78	2.27	.02	.74	.88	.13	.84	.45	.57	(ь)				
AL,03	31.10	3. 79	1.86	1.93	33.00	4.28	2.51	32.05	2.76	2.56	2.66	31.25				
Cr,03	(Ь)	.81	. 83	.82	(Ъ)	(b)	(b)	(ь)	(b)	.06	. 05	(b)				
FeU	.49	9.82	14.20	22.18	. 45	14.45	19.30	. 59	13.70	15.00	19.87	.45				
MnD	(Ъ)	.17	.26	.02	(b)	(b)	(b)	(Ъ)	(b)	.21	. 22	(в)				
Mg0	. 18	29.43	25.86	8.01	. 16	24.72	21.20	.11	25.51	23.81	19.98	.00				
Ca0	:6.48	1.41	2.39	16.43	15.38	2.76	2.89	16.21	2.21	2.11	2.82	16.06				
Na ₂ 0	1.70	(b)	(b)	(ь)	1.70	.06	.03	1.51	.03	(Ъ)	(b)	1.58				
к ₂ 0	.22	(b)	(b)	(b)	.18	.01	.02	. 29	.01	(b)	(b)	. 20				
Total	98.24	99.80	99.02	99.64	99.51	99.46	99.66	99.20	98.49	99.39	99 .30	98.90				
					Unit	formulas										
Si	2.242	1.901	1.930	1.907	2.230	1:901	1.95	2.232	1.945	1.994	1.975	2.280				
Ti	.004	.016	.022	.068	.001	.021	.02	.005	.023	.013	.016					
A1 ^{IV}	1.714	.099	.060	.091	1.784	.099	.04	1.746	.055	.066	. 025	1.702				
41 ^{VI}		. 059	.021	.000	د.	.084	.06		.064	. 104	. 092	••				
Cr		.023	.024	.003		- •		••	••	.002	. 002					
Fe	.019	.291	.434	.738	.018	.439	.59	.024	.418	.454	. 618	.018				
Mn		.006	.009	.001						.007	. 007					
Mg	.013	1.550	1.408	.475	.012	1.336	1.16	.008	1.384	1.282	1.107	••				
Ca	.826	.054	.0%	.700	.756	.108	.11	.803	.087	.082	.113	.795				
Sa	.155	••	••		. 152	.005	.00	.135	.003			. 142				
К	.014				.011	.001	.00	.018	. 002			- 012				
					Atom	ic ratios										
	Ca 83.0	Mg 81.8	Mg 72.7	Mg 24.8	Ca 82.3	Mg 70.9	Mg 62.1	Ca 84.0	Mg 73.3	Mg 70.5	Mg 60.2	Ca 83.8				
	Na 15.6	Fe 15.4	Fe 22.4	F & 38.8	Na 16.5	Fe 23.3	Fe 31.7	Na 14.1	Fe 22.1	Fe 23.0	Fe 33.6	Na 15.0				
	K 1.4	Ca 2.8	Ca 4.9	Ca 36.6	K 1.2	Ca 5.8	Ca 6.2	K 1.9	Ca 4.6	Ca 4.5	Ca 6.2	K 1.2				

^aFrom table 1-V(a).

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1.14

^bNot determined.

THE HISTORY OF LUNAR BRECCIA 15015

⁽b) Continued

TABLE 1-V.- Concluded

(b) Concluded

Compound			_		Analysis no.	, ^a wt.7					
or element	37	38	39	40	41	42	43	44	45	46	47
sio ₂	97.99	••	37.84	53.66	45.45	38.20	53.45	44.10	50.51	44.66	53.17
TiO,	(6)	51.03	.82	.96	.27	. 18	.74	.00	2.00	.00	.76
A1,0,	1.70	1.84	.05	.80	34.06	.76	2.36	34.54	. 92	35.85	1.71
Cr.0.	(Ъ)	. 14	(ь)	(ь)	(5)	(b)	(b)	(ь)	. 29	(ь)	(b)
FeU MnO MgO CaO Na ₂ O K ₂ O	.27 (b) .00 .02 (b) (b)	46.79 .10 .19 (b) (b) (b)	26.84 (b) 35.06 .15 .02 .02	16.30 (b) 25.78 1.61 .03 .02	.39 (b) .00 17.58 1.20 .19	22.89 (b) 37.33 .01 (b) (b)	13.67 (b) 26.50 1.68 .22 .01	.13 (b) .13 18.78 46 .06	10.03 .22 12.83 20.77 (b) (b)	.32 (b) .02 18.79 .77 .11	14.81 (b) 26.66 1.80 .00
Total	99.98	100.09	100.80	99.16	99.14	99.37	98.63	98.40	97.57	100.52	98.92
		*		·	Unit form	ulas					•
Si Ti		0,963	0.997 .017	1.964	2.115	1,000 .001	1.942	2.071	1.944 .058	2.054	1.957
AI AI ^{VI} Cr		.055 .003				.024	.058		.042	1.944 	. 04 3
Fe		. 981	.591	. 499	.016	.502	.416	.006	. 323	.013	.456
42 MD		.003 .006	1.376	1.406	.000	1.457	1.435	.010	.008 .736	.002	1.408
Ca			.005	.063	.877	.001	.066	.945	.857	. 927	. 071
K K			.001	.003	.109		.016 .001	.061 .004		.069	.000
		L		I	Atomic ra	 tios					L
		₩: 0.8 Fe 99.2	Mg 70.0 Fe 30.0	Mg 71.4 Fe 25.4 Ca 3.2	Ca87.9 Na10.9 K 1.2	Mg 74.4 Fe 25.6	Mg 74.9 Fe 21.7	Ca 93.6 Na 6.0 K .4	Mg 38.3 Fe 17.2 Ca 44.5	Ca92.4 Na 6.9 K .7	Mg 72.7 Fe 23.6 Ca 3.7

^aFrom table 2-V(a).

^bNot determined.

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LUNAR SAMPLE STUDIES

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THE HISTORY OF LUNAR BRECCIÁ 15015

are possibly recrystallized rocks in which members of the Fra Maturo or KREEP suite or their fragmental associates predominate: e.g., analyses 39 to 43, table $1 \cdot V(b)$. In both groups, homogeneous hypersthene is the dominant ferromagnesiant phase. The Fe/Ee+Mg ratio of approximately 0.25 in the hypersthene is within the range of zoned hypersthene in the Fra Mauro basalts. The calcium (Ca) content is marginally lower and the aluminum (Al) content distinctly less in the metamorphic pyroxenes.

No fragments of rocks similar to members of the mare basalt group with a metamorphic texture have been found in 15015. The same is true for the feld-spathic basalt group. Reference to figure 1-8 shows an absence of metamorphic pyroxenes with the Fe/Fe+Mg ratios >0.3 that are characteristic of these igneous groups, which reinforces the tentative conclusion that the feldspathic basalts are variants of the mare group of basalts. These basalts should be grouped together as younger effusives that postdated the inspact events that were ultimately responsible for the metamorphic nature of a proportion of the clasts here assigned to the Fra Mauro and ANT suite.

A Tentative History of 15015 Based on General Petrology

Breecias 15015 and 15021, nearly identical in chemical composition, were collected on a ray from Aristillus or Autolycus just west of the LM and near the castern limit of the mare surface overlooked by the mountains of the Apennine front (ref. 1-19). This suggests that the rock 15015 could be a boulder from a distant source, which lay partially buried in the finer grained, Jess coherent ray material, or that it was buried and /reexcavated by some minor cratering event. Altornatively, this boulder may have been formed by shock-lithification of unconsolidated ray material/by some local cratering event that deposited it where it was collected. In the first case, lithification was effected by or preceded by the Aristillus event, and the material of the boulder reflects only the lithology of its distant source. In the second and more probable case, during the interval between ray formafion and shock lithification, material of more local origin could have been mixed with the surface layers of the ray material and formed an additional component of the shock-lithified ejecta. Reid et al. (ref. 1-12) have shown that 15021 is enriched in Fra Mauro basalt glass, and Powell et al. (ref. 1-20) have shown that Fra Mauro basalts are more abundant in the <4-mm lithic fragments at the LM site than at other mare sites visited during the Apollo 15 mission. The same features are found in the vitric and lithic clasts of 15015 and are considered diagnostic of the ray. Therefore, the rocks from which these clasts were der 1 possibly underlie the mare basalts of the Autolycus-Aristillus region. The overlying spectrum of mare basalts should also be found in the ray material. Possibly the feldspathic basalts of 15015, which were assigned previously to the mare basalt group, are the representatives of mare basalts of this distant region.

The smail amount of metamorphic rocks in 15015 could be of local (Apennine front) origin or could have come from a distant source: a major crater like Aristillus could be expected to excavate fallback material from the Imbrium event.

The green glass spheres are tound as a subordinate component of the glasses in the <1-mm fraction (15021) of the ray soil. They are also a subordinate component in the shock-lithified boulder of soil breccia 15015, but here they are all in a devitrified state. The wide distribution of the green glass spheres and their greater concentration in many of the <1-mm fractions of the surface regolith samples from the Apollo 15 site suggests that they are not a part of the original ray material, but that they were deposited after ray formation and before the cratering event that shock-lithified 15015 and deposited it as a surface boulder.

Selenchronology

Six samples of 15015,15 have been subjected to ${}^{40}\text{Ar}{}^{39}\text{At}$ analysis. They were: 5c, a 60-mg sample of the dark matrix containing numerous small inclusions; 5b, fragments (33 mg) of a large gray clast chipped from the dark matrix; 23c (52 mg), 23d (19 mg), 26, fragments (24 mg) of bubbly glass chipped from the upper surface of the rock; and 23b, fragments (20 mg) of a white clast embedded in the surface of rock 15015 and partially included in the bubbly glass. The mineralogy of the rock fragments has been described in detail previously. Sample 5b is a dark variolitic basalt with >30 percent plagioclase (feldspathic basalt). Sample 23b is a pale mesostasisrich basalt (Fra Mauro type). The matrix and glass

were not subjected to a separate mineralogical analysis, though analyses 1 and 2 (table ¹-II), respectively, can probably be taken as representative. Details of the irradiation (designated SII15) have been published elsewhere (ref. 1-6).

Each sample was subjected to a stepwise heating experiment, and the Ar-release patterns obtained are summarized in table 1-VI. Apparent K-Ar ages have been calculated from the gas released at each temperature step for the two clasts and are shown in figs. 1-10 and 1-11. The amount of cosmogenic ${}^{38}\text{Ar}_{c}$ is calculated for each temperature assuming a binary



Figure 1-10.- Argon isotope release pattern for Fra Mauro basalt-type fragment 23b, indicating a 75-percent loss of radiogenic ${}^{40}Ar$ and a 20-percent loss of cosmogenic ${}^{38}Ar_c$. The high-temperature ${}^{40}Ar/{}^{39}Ar$ age attains a value of 3.7 \pm 0.1 b.y. despite the extreme loss and is comparable to, but slightly lower than, more precise ages of other Fra Mauro basalts. A well-defined cosmic ray exposure age of 490 m.y. is determined.



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THE HISTORY OF LUNAR BRECCIA 15015

mixture of cosmogenic Ar $({}^{36}\text{Ar}/{}^{38}\text{Ar} = 0.65)$ and trapped Ar $({}^{36}\text{Ar}/{}^{38}\text{At} = 5.35)$. The ratio ${}^{38}\text{Ar}_c/$ ${}^{37}\text{Ar}$ is plotted also in figures 1-10 and 1-11 and is related directly to the ${}^{38}\text{Ar}_c/\text{Ca}$ ratio and hence to the cosmic ray exposure age (ref. 1-21).

Also plotted in figures 1.10 and 1.11 is the ratio 39 Ar*/ 37 Ar, which is proportional to the average K/Ca ratio of the sites releasing Ar at each-stage of the experiment². The overall K/Ca ratio for the

 240 Ar⁺ and 39 Ar⁺ refer to Ar produced by radioactive decay of 40 K and (n,p) reactions on 39K, respectively.

matrix, 0.024, is identical to that of other matrix samples (analyses 14 and 15, table 1 II) and is slightly smaller than that of the glass, 0.030. The K/Ca ratio of the gray clast (5b) is much lower, 0.0051, while that of the white clast (23b) is high 0.062, reflecting the presence of a KREEP component.

Both clasts show the effects of extreme radiogenic 40 Ar loss, at least 75 percent, based on the maximum 40 Ar $^*/{}^{39}$ At * ratios reached in the high-tentperature release. Variations in the 39 Ar $^*/{}^{37}$ Ar ratio indicate that the loss is associated with low-retentivity, potassium-rich sites. A parallel loss of cosmogenic

TABLE 1-VI	BRECCIA	15015,15	ARCON-RELEASE	PATTERNS
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Temperature, K (°C)	³⁶ Ar/ ³⁸ Ar	³⁸ Ar/ ³⁷ Ar	³⁸ Ar _c / ³⁷ Ar (a)	³⁹ Ar*/ ³⁷ Ar	⁴⁰ Ar/ ³⁹ Ar*	³⁹ Ar [#] (b)	Apparent age, b.y. (c)
		1501	5,15,23b (white cl	aat, 33.0 mg); J = 0	.02982		
773 (500) 873 (600) 973 (700) 1043 (770) 1123 (850) 1193 (920) 1303 (1030) 1403 (1130) 1573 (1300) Total	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	0.0345 ± 11 .0180 ± 9 .0138 ± 4 .0303 ± 3 .0543 ± 3 .0633 ± 4 .0669 ± 4 .0580 ± 5 .0571 ± 6 .046	 0.0134 ± 4 .0285 ± 3 .0474 ± 3 .0563 ± 4 .0558 ± 4 .0537 ± 5 .0553 ± 6 .042	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	18.6 28.1 18.0 13.6 6.1 3.7 2.9 .8 1.2 90	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
			015,15,5b (gray cla	ast, 20.0 mg); J = 0	. 02982		
773 (500) 873 (600) 973 (700) 1023 (750) 1073 (800) 1123 (850) 1133 (920) 1343 (1070) 1573 (1300) Total	1.14 ± 0.30 .87 ± .21 .90 ± .02 1.06 ± .01 1.60 ± .01 1.42 ± .01 1.28 ± .01 1.40 ± .01 1.28	0.0211 ± 37 .0096 ± 19 .0177 ± 4 .0624 ± 5 .1380 ± 12 .1720 ± 2 .1714 ± 6 .1719 ± 5 .1629 ± 1 .137	0.0189 ± 40 .0092 ± 20 .0168 ± 4 .0569 ± 5 .129 ± 1 .159 + .143 ± 1 .149 + 1 .137 ± 1 .119	0.221 ± 12 .092 ± 6 .0253 ± 18 .0055 ± 14 .0067 ± 20 .0102 ± 25 .0055 ± 18 .0018 ± 8 .0011 ± 3 .0093	0.3 ± 0.3 1.0 ± .3 10.4 ± .9 60.9 ± 7.0 146 ± 40 156 ± 40 156 ± 60 180 ± 80 203 ± 50 49.5 • 0.0298	2.0 2.4 1.9 1.1 .58 .55 .39 .23 .57 9.8	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
	F	[]	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	B10851 2112	- 0.0298		
773 (500) 873 (600) 973 (700) 1043 (770) 1123 (850) 1193 (920) 1343 (1070) 1573 (1300) 1743 (1470) 7.41	5, 11 + 0.09' $5.32 + .06$ $5.32 + .02$ $5.33 + .04$ $5.33 + .03$ $5.31 + .03$ $5.11 + .02$ $5.11 + .04$ $5.14 + .01$ 5.11	$1.23 + 0.22$ $.64 + .04$ $.88 \pm .02$ $1.61 + .03$ $2.91 \pm .03$ $3.46 + .04$ $3.73 \pm .02$ $1.56 \pm .01$ $1.80 + .01$		0.352 ± 63 .255 ± 16 .157 ± 4 .113 ± 2 .099 ± 2 .0504 ± 5 .0367 ± 6 .0462 ± 2	$\begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$	0.4 1.3 2.5 2.7 4.4 3.2 9.3 9.3 9.2 16.7	⁴ 2.08 ± 0.04 ⁴ 1.18 ⋅ .03 ⁴ 1.25 ⋅ .02 ⁴ 1.29 ⋅ .04 ⁵ 1.21 ⋅ .08
Total	5.17	2.22		,0549	273.4	69.7	

^aThe ³⁸Ar_c/³⁷Ar rdtio was calculated assuming that ³⁸Ar originaton solely from common Ar ((³⁶Ar/³⁸Ar)_c = 0.65) and trapped Ar ((³⁶Ar/³⁸Ar)_t = 5.35). The corresponding ³⁸Ar_c/Co ratio is given by ³⁸Ar_c/Co = 4.14 × 10⁻³ × J × (³⁸Ar/³⁷Ar) cm³ standard temperature pressure (STP)/gm Co, where J is the irradiation coefficient (ref. 1-21).

^bAmounts in units of 10^{°8} cm³ STP/g. Uncertainty in absolute amounts is *20 percent.

 $^{\circ}$ = 5.305 = 10⁻¹⁰ v⁻¹, $^{\circ}$ = 0.585 = 10⁻¹⁰v⁻¹, 40 K/K = 0.0119 percent.

 $1 = (exp(t) = 1)/40 Ar/39 Ar^{+})$. No correction applied for trapped/cosmogenic 40 Ar except under (d),

dApparent ages calculated assuming (⁴⁰Ar/³⁰Ar), = 0.97 for trapped component in Sc and (⁴⁰Ar/¹⁶Ar), = 1.15 for 23c, .3d and 26.

LUNAR SAMPLE STUDIËS

TABLE 1-VI. - Concluded

Temperature, K (°C)	³⁶ Ar/ ³⁸ Ar	³⁸ Ar/ ³⁷ Ar	³⁸ Ar _e / ³⁷ Ar (a)	³⁹ Ar*/ ³⁷ Ar	⁴⁰ ar/ ³⁹ ar#	³⁹ Ar# (h)	Apperent age, b.y. (c)
· · · · · · · · · · · · · · · · · · ·		1501	3,15,23d (voulcula	or 81665, 19 m8); J •	0.0256		
803 (530)	5.37 ± 0.15	0.903 : 26	••	0.134 ± 58	720.0 ± 320,0	0.1	••
903 (630)	5.27 t .10	.830 ± 18	••	.186 ± 18	157.0 ± 16.0	.7	d _{2,8} ± 0,2
993 (720)	5,26 1.03	1.046 ± 8		.136 2 5	112.2 ± 4.1	2.2	^d 1.9 2 .1
1093 (820)	5.24 t .03	1,096 ± 7		.0979 ± 13	101.0 ± 1.5	4,3	^d 1.2 ± .1
1173 (900)	5.28 ± .04	1,600 ± 15	••	.0852 ± 14	170.1 ± 2.9	3.2	^d 1.7 ± .1
1263 (990)	5.20 ± .01	1.847 ± 12	••	.0736 ± 21	225.2 ± 6.3	2.8	d _{2.0 t} .2
1353 (1080)	1.16 ± .04	2.841 ± 23	••	.0608 ± 4	305.0 2 1.4	11.8	••
1473 (1200)	4.81 + .01	.776 ± 4		.0275 + 12	270.0 + 12.0	12.9	
1583 (1310)	4.82 ± .08	.653 ± 14	••	.0230 ± 71	840.0 ± 260.0	.2	
1723 (1450)	4.77 ± .06	.568 ± 13	••	.0254 ± 62	660.0 ± 170.0	.2	••
Tot al	5.16	1.70		0.054	233.2	40.0	••
		1501	5,15,26 (vesicula	r glass, 24 mg); J =	0.0256		
803 (530)	5.29 ± 0.11	1.034 * 35		0.239 17	102.8 ± 7.9	0.7	^d 2.0 ± 0.2
903 (630)	5.35 ± .06	.682 11		.138 ± 7	50.2 ± 3.2	1.3	d,77 ± .10
993 (720)	5.34 ± .03	1.315 ± 19		.111 ± 3	94.4 ± 2.5	2.1	^d .88 ± .08
1073 (800)	5.25 ± .04	2.467 : 26		.0962 ± 18	181.6 ± 3.1	3.4	Bo. + ee.b
1173 (900)	5.20 ± .01	3,000 ± 19		.0819 ± 17	255.9 ± 5.1	2.6	^d 1.3 ± .2
1263 (990)	5.15 ± .03	2.476 ± 21	••	.0500 ± 20	329.6 ± 12.8	1.9	d _{1.2} , 3
1353 (1080)	5.20 ± .03	4.353 ± 28		.0472 ± 3	553.0 ± 2.7	9.1	••
1413 (1140)	5.14 ± .03	2.279 ± 18	••	.0308 ± 9	454.5 ± 12.4	4,1	••
1723 (1450)	4.91 2 .05	1.068 ± 19		.00%8 17	590.0 2 120.0	.2	
Total	5, 18	3.04		0.052	365.2	25.6	
			15015.15.5c (mate	ix. 60 mg); J = 0.029	82		<u> </u>
773 (500)	5.25 ± 0.08	0.74 ± 0.15		0.45 . ± 0.09	31.3 ± 0.9	1.6	^d 0,98 ± 0.03
873 (600)	5.28 ± .05	.45 ± .03	••	,281 ± .019	15.1 ± .2	2.9	^d .35 ± .01
973 (700)	5.31 ± .03	.58 ± .01		.153 ± ,003	31.4 1 .3	4.3	^d ,57 ± .01
1043 (770)	5.27 ± .03	2.31 ± .03	••	.086 ± .002	162.8 ± 2.6	3.9	d1,06 ± .02
1123 (850)	5.22 ± .02	4.94 ± .07	••	.060 ± .001	444.7 ± 7.2	2.8	••
1193 (920)	5.25 2 .02	0.08 ± .15		.055 ± .002	555.6 * 8.4	2.4	••
1343 (107C)	5.25 ± .02	8.71 ± .28		.034 2 .002	85 2 910	11.7	
1393 (1120)	5.19 ± .02	4.13 2 .05	••	.022 + .001	950 24	1.1	••
1473 (1200)	5.07 ± .03	1.62 ± .04	••	.014 .001	662 ± 46	1.0	••
1553 (1280) 1743 (1470)	5.19 ± .04 4.89 + .05	2.18 ± .07		.019 ± .001	587 ± 31	.5	
		1,46 I ,4J			"	.02	
Total	5.22	5.0		0.044		40.6	

^aThe ³⁸Ar_c/³⁷Ar ratio was calculated assuming that ³⁸Ar originates solely from cosmogenic Ar $(({}^{36}\text{Ar}/{}^{38}\text{Ar})_c = 0.65)$ and trapped Ar $(({}^{36}\text{Ar}/{}^{38}\text{Ar})_t = 5.35)$. The corresponding ${}^{38}\text{Ar}_c/\text{Ca}$ ratio is given by ${}^{38}\text{Ar}_c/\text{Ca} = 4.14 \times 10^{-3} \times \text{J} \times ({}^{38}\text{Ar}/{}^{37}\text{Ar})$ cm³ standard temperature pressure (STP)/gm Ca, where J is the irradiation coefficient (ref. 1-21).

^bAmounts in units of 10⁻⁸ cm³ STP/g. Uncertainty in absolute amounts is +20 percent.

 $c_{\lambda} = 5.305 \times 10^{-10} \text{ y}^{-1}, \lambda_{\mu} = 0.585 \times 10^{-10} \text{ y}^{-1}, \frac{40}{8} \text{ K/K} = 0.0119 \text{ percent}.$

J = $(e_{\pi p}(\lambda t) - 1)/{^{40}Ar}/{^{39}Ar^{40}}$. No correction applied for trapped/cosmogenic ${^{40}Ar}$ except under (d).

^dApparent ages calculated assuming (40 Ar/ 36 Ar), = 0.97 for trapped component i... ic and (40 Ar/ 36 Ar), = 1.1° for 23c, 23d and 26.

 ${}^{38}\text{Ar}_{c}$ is in the low-temperature release, evidenced by low ${}^{38}\text{Ar}_{c}/{}^{37}\text{Ar}$ ratios and suggesting that some of the Ar loss occurred after the main period of cosmic ray exposure of the fragments. The chemical similarity of glass and matrix (section entitled "Mineralogy and Petrology") suggests that the glass was formed by melting and lithification of a preexisting soil. Such an event could certainly account for the extreme Ar loss in this sample. However, it should be noted that 50 percent of the artificially induced 39 Ar* was released by 873 K (600° C) in both clasts, and it therefore seems quite possible (ref. 1-22) that

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some of the loss could be accounted for by solar heating.

In the last 15 p, reent of ${}^{39}Ar^*$ release from 23b, the ${}^{40}Ar^*/{}^{39}Ar^*$ ratio reaches a mean value corresponding to an apparent age of 3.71 ± 0.12 b.y. This must be regarded as a lower limit to the original crystallization age. The high-temperature release from 5b (1073 to 1573 K (800° to 1300° C)) corresponds to an ${}^{40}Ar^{-39}Ar$ age of 3.4 ± 0.2 b.y. Because of the extreme degree of ${}^{40}Ar^*$ loss and the low K content of 5b, this age is rather inaccurate and again must be regarded as a lower limit to the original crystallization age.

The major release of cosmogenic Ar occurs at high temperature, and well-defined, but distinct, ${}^{38}\text{Ar}_c/{}^{37}\text{Ar}$ plateaus are obtained for both clasts. Based on a nominal ${}^{38}\text{Ar}_c$ production rate of 1.4×10^{-8} cm³ standard temperature pressure (STP)/g Ca/m.y. (ref. 1-21), exposure ages of 490 m.y. (23b) and 1290 m.y. (5b) are calculated. Ages based on total ${}^{38}\text{Ar}_c/{}^{37}\text{Ar}$ are respectively 400 m.y. and 1050 m.y. and reflect some 20 percent ${}^{38}\text{Ar}_c$, loss.

The Ar data for the matrix (5c) and the glasses (23c, 23d, 26) are dominated by trapped Ar, and age information is less clear. A graph of ⁴⁰Ar/³⁶Ar against ³⁹Ar*/³⁶Ar (fig. 1-12) reveals an apparent correlation for glasses 23c and 26, corresponding to isochron ages of 1.2 b.y. and 0.8 b.y. The correlations are largely defined by relatively low temperature points (<9273 K (9000° C)) and are subject to numerous interpretations. Sample 23d shows no correlation. The frothy glass contains many small lithic and mineral inclusions, and it is suspected that the lowtemperature release, being relatively low in trapped Ar, may be dominated by these inclusions while the glass itself may be responsible for the greater proportion of trapped Ar in the high-temperature Ar. It seems likely that the isochron ages may represent at best an upper limit to the age of the glass, on the assumption that the lithic clasts were largely outgassed at the time the glass was melted. The ⁴⁰Ar/ ³⁶Ar ratio of the trapped Ar in the glass is $1.17 \pm$ 0.04, as indicated by the intercept in figure 1-12. This is 0.20 ± 0.06 , higher than the corresponding ratio for the matrix. The difference could have resulted from the homogenization of trapped Ar and radiogenic Ar within the glass at the time of its formation and to



Figure 1.12.- Isochron diagram of ${}^{40}\Lambda r/{}^{36}\Lambda r$ and ${}^{39}\Lambda r^{\circ}/{}^{36}\Lambda r$ for Λt from vesicular glasses 23c. 23d, and 26. The numbers represent temperature in units of 100° C ($T_{K} = T_{C} + 273.15$). The slopes of the isochron for 23c and 26 correspond to ages of 1.2 and 0.8 b.y., respectively. No correlation is observed for 23d, and the dashed line drawn corresponds to the same apparent age as 23c. The correlations appear to indicate outgassing of the glass or of inclusions within the glass within the last 1.0 h.y. It has been tentatively inferred that the glass formed at 1.0 b.y. or later, but the correlations could reflect recent solar heating. (a) Sample 15015.23c. (b) Sample 15015.23d. (c) Sample 25025.26.

this extent is consistent with the glass being relatively recent.

The matrix sample (5c) does not yield a welldefined correlation of ${}^{40}\Lambda r/{}^{36}\Lambda r$ and ${}^{39}\Lambda r^*/{}^{36}\Lambda r$. Above 1223 K (950° C), the yield is entirely dominated by trapped Ar with a 40 Ar/ 36 Ar ratio of 0.97 \pm 0.04. Apparent ages for the low-temperature release steps are calculated assuming a trapped component of this composition and are included in table 1-VI. The apparent ages are low, reflecting Ar loss, as observed for the clasts and possibly for the glass.

The predominance of trapped Ar in the matrix and glass precludes an accurate estimation of an ³⁸Ar exposure age for these samples. The ³⁶Ar/³⁸Ar-ratio (table 1-VI) decreases by a few_percent in the hightemperature release, which is consistent with a small cosmogenic contribution. An ${}^{38}\text{Ar}/{}^{36}\text{Ar}$ and ${}^{37}\text{Ar}/{}^{36}\text{Ar}$ correlation diagram (fig. 1-13) appears to indicate the absence of cosmogenic ³⁸Ar in the lowtemperature release (approximately 10 percent of total ³⁷Ar release). The ³⁸Ar/³⁶Ar ratio in the hightemperature telease increases with increasing ³⁷Ar/ ³⁶ Ar in a fashion approximately consistent with the presence of cosmogenic ³⁸Ar. The data for both matrix and glass do not appear to be in conflict with an exposure age similar to that of 23b, but the total variation in ${}^{38}i$ r/ ${}^{36}Ar$ is only a few percent. In view of the possibility of fractionation effects at this level due to thermal diffusion, it is not possible to make any precise comments on the exposure age data of matrix and glass.

The results of these measurements provide a number of constraints for models of the possible history of 15015, but they do not define a unique history. The constraints are as follows.

1. The high-temperature ${}^{40}\text{Ar}*/{}^{39}\text{Ar}*$ ratios from the Fra Mauro basalt clast (23b) indicate that the parent rock crystallized at 3.7 ± 0.1 b.y. or earlier, predating the Hadley Rille mare basalts (refs. 1-23 to 1-26).

2. The variolitic basalt (5b) crystallized at 3.4 ± 0.2 b.y. or earlier.

3. Although no mare basalt clasts from 15015 have been dated, their presence (section entitled "Mineralogy and Petrology") indicates that the breccia formed after the basalt flows had crystallized. If the mare basalt fragments are locally derived, this would imply that the breccia formed more recently than 3.3 b.y. ago (refs. 1-23 to 1-26).

4. The variolitic basalt clast (5b) has an exposure age of 1290 m.y., greatly in excess of that of the Fra Mauro basalt clast 23b, 490 m.y., and also greatly in excess of neon-21 (21 Ne) exposure ages inferred for the matrix (noble gas data). This implies a preirradia-



Figure 1.13.- Correlation diagram of ${}^{38}\text{Ar}/{}^{36}\text{Ar}$ and ${}^{37}\text{Ar}/{}^{36}\text{Ar}$ for matrix 5c and glasses 23c, 23d, and 26. The Ar is a mixture of trapped Ar (${}^{38}\text{Ar}/{}^{36}\text{Ar} = 0.188$, ${}^{37}\text{Ar}/{}^{36}\text{Ar} = 0$) and cosmogenic Ar produced predominantly from the bombardment of Ca (${}^{38}\text{Ar}/{}^{36}\text{Ar} \approx 1.6$, ${}^{37}\text{Ar}/{}^{36}\text{Ar} \neq 0$, inversely proportional to exposure age). For all samples, the low-temperature release lies on a horizontal line, reflecting the absence in these (Ca) sites of cosmogenic ${}^{38}\text{Ar}_c$ and ${}^{36}\text{Ar}$. The higher temperature points show the release of cosmogenic ${}^{38}\text{Ar}_c$ and ${}^{36}\text{Ar}$ in amounts approximately proportional to the amount of ${}^{37}\text{Ar}$ (= Ca). The upper line corresponds to an exposure age of 1000 m.y. irradiation coefficient (J = 0.0298) or-1160 m.y. (J = 0.0256). The slope of the middle line corresponds to an exposure age of half this. The absence of cosmogenic Ar in the low-temperature points appears to be at odds with the observations of radiogenic ${}^{40}\text{Ar}$ (fig. 1-12).

tion of clast 5b for at least 800 m.y. before incorporation in the breccia. Such a history of preirradiation combined with the ${}^{40}\text{Ar}*/{}^{39}\text{Ar}*$ age indicates that 15015 was formed more recently than 2.6 ± 0.2 b.y. ago from a mature regolith. Petrologic considerations (discussed earlier) suggest that this regolith was locally derived. In view of the late date of the lithification in comparison to the time of basalt extrusion, it appears likely that lithification resulted from meteorite impact.

5. The 40 Ar 39 Ar isochron ages of 23c and 26, around 1.0 b.y., may represent the formation time of

24

THE HISTORY OF LUNAR BRECCIA 15015

the glass and also the time of the lithification, although this interpretation is speculative. If the glass does date from this time, it is apparent from the track and ³He measurements (discussed in following sections) that the sample remained buried at depth until quite recently.

Cosmic Ray Tracks

Combined high-voltage and scanning electron microscopy, previously applied to Apollo 11 and 14 breccias (refs. 1-27 and 1-28) have been used to study both radiation damage features and various textural features in individual grains from breccia 15015.

Preliminary 1-MeV electron microscope examination of micron-sized grains extracted from the bottom chip, 15015,15,10, yields the following results. 1. Less than 10 percent of the grains appear as totally amorphous, but a much higher proportion shows evidence of shock (i.e., diffraction patterns indicative of disorder and deformation bands in _ pyroxenes).

2. No evidence exists for ultrathin amorphous coatings or high densities of latent tracks characteristic of highly irradiated particles in mature. soil samples.

3. Approximately 30 percent of the grains contain microcrystallites (fig. 1-14) similar to those found both in artificially heated lunar (ust grains and in mildly metamorphosed Apollo 11 and <u>-14 breccias.</u> belonging to groups 1 and 2 (ref. 1-29).

Polished sections from two exterior chips, 15015,15,10 (bottom) and 15015,15,22 (top), have been irradiated with a californium (^{252}Cf) source to induce fission-fragment tracks and then examined by



Figure 1-14.- 1-MeV electron micrographs of crystalline grains loaded with microcrystallites. (a) Dark field. (b) Bright field.

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LUNAR SAMPLE STUDIES

an SEM. Gradual etching was used to reveal both artifical and fossil tracks successively in glass, feldspars, and pyroxenes (each mineral phase being identified with the electron microprobe attachment of the SEM). The main results are as follows.

1. A very slight etching quickly dissolves the finegrained matrix that presumably consists of glassy or highly disordered grains. Glass is also found in vesicular veins and in numerous angular clasts, and constitutes the external coatings of the two chips. All interior glassy grains have fossil track densities of approximately 10⁶ tracks/cm². It is almost impossible to use the exterior glass coating as a track detector because high concentrations of vesicular bubbles are superimposed on the track distribution. Therefore, no track density gradient can be identified, and an approximate estimate indicates the density ρ is of the order of 10⁷ tracks/cm². The very heavy exposure age of the glass coating is therefore <10⁷ years.

2. Feldspar grains are very scarce, very small (<10

 μ m), and surrounded by a much larger area of disordered mineral phase of anorthosite composition (fig. 1-15). Eleven such grains have been found with track densities at the centers ranging from 10⁷ to 4 X 10⁹ tracks/cm².

3. Pyroxene grains are highly fractured and have track densities at the center ranging from about 10^6 to 4×10^8 tracks/cm².

In a previous study of Apollo 11 and 14 breccias, two distinct groups of breccias were defined in terms of track metamorphism (refs. 1-27_and 1-28) as follows.

1. The mildy metamorphosed breccias still contain radiation damage features due to the individual irradiation of their constituent grains in solar nuclear particle fluxes before brecciation. Such features appear either as high densities of latent nuclear particle tracks (approximately 10^{11} tracks/cm²) or as very small crystallites similar in size to those formed. in artificially heated mature lunar dust grains. These



Figure 1-15.- Scanning electron micrographs of a polished section that has been etched to reveal tracks in feldspars. (a) Aluminum X-ray scan superimposed on a backscattered electron scan. The arrow indicates the location of a feldspar grain of high track density in a rich Al-bearing mineral phase. (b) Micrograph of the same feldspar grain of high track density.

breccias are also characterized by high densities (approximately 10^9 tracks/cm²) of etched tracks at the center of 100- μ m grains.

2. In the stronglv metamorphosed breccias, no remnant of the prebreccia solar irradiation is found, and the track densities of about 10^7 tracks/cm² seem to be due to the exposure of the whole breccia to galactic cosmic rays.

Rock 15015 is intermediate between these two extreme groups. It contains microcrystallites similar to those observed in mildly metamorphosed breccias but much larger in size. Furthermore, the track densities in feldspars and pyroxenes are spread over two orders of magnitude, in striking contrast to the narrow range of values observed for the second group of breccias. These "intermediate" characteristics suggest either that the breccia has been formed from a mixture ofmature and nonmature soil or that a nonthermal process has also been responsible for the track metamorphism. At present, the second explanation is preferred, because several textural features (extensive fracturing, lattice disorder) support the hypothesis that rock 15015 has suffered a strong shock event that could have severely affected its track distribution.

Noble Gases

Concentrations and isotopic compositions of He, Ne, and Ar were determined by mass spectrometry of three matrix samples and of one clast sample of 15015,15 (table 1-VII).

The spallogenic and trapped Ne components are evaluated by assuming a ${}^{22}Ne/{}^{21}Ne$ ratio of $1.10 \pm$ 0.05 in spallogenic Ne and one of 32 ± 1 in trapped Ne. These calculated concentrations are given in table 1-VII. Together with the chemical composition given in table 1-II and respective elemental production rates, the ${}^{21}Ne$ exposure ages are computed (table 1-VII). Within limits of error, the ${}^{21}Ne$ exposure ages of the three matrix samples and the clast sample agree with each other. Unfortunately, the chemical composlition of the clast was not determined, which renders the ${}^{21}Ne$ exposure age uncertain despite the comparatively small error of the spallogenic ${}^{21}Ne$ component.

The predominance of trapped Ar renders the calculation of the spallogenic Ar components uncertain,

except for the clast sample. Assuming an ³⁶At/³⁸Ar ratio of 5.35 ± 0.05 for trapped Ar and one of $0.65 \pm$ 0.03 for spallogenic Ar, a concentration of spallogenic ³⁸Ar approximately 30×10^{-8} cm³ STP/g is calculated for the clast. Spallogenic ³⁸Ar is also clearly present in the bottom and top saniples, but the quantities are uncertain. With a Ca concentration in the clast assumed to be equal to that in the matrix, an exposure age of 290 \pm 20 m.y. is determined. Using the formula for the production rate of 38 Ar. given by reference 1-31, an exposure age of 203 ± 56 m.y. is obtained. An evaluation of the radiogenic and trapped ⁴⁰Ar components is not possible. As seen from the measured 40 Ar/36 Ar ratios, a single 40 Ar/36 Ar 36 Ar ratio in the Ar initially trapped and a constant concentration of radiogenic ⁴⁰Ar can both be ruled out. This is presumably due to an inhomogeneous K distribution. With the K concentration given in table 1-II, approximately 10^{-4} cm³ STP/g of radiogenic ⁴⁰Ar would be expected if the sample had retained Ar for 4 b.y. Clearly, the clast sample has either retained Ar for a shorter period or is depleted in K.

Concerning the composition of the He in the 15015 samples, the following observations are made. The 4 He/ 3 He ratios of trapped He in lunar fines are around 2800, and the 3 He/ 21 Ne ratio of spallogenic gas, around 6. Thus, a substantial deficit of 3 He is evident in all samples. Also, the measured 4 He/ 20 Ne ratios are low compared to lunar soil values of approximately 20 to 50. This indicates also a substantial deficit of 4 He. Taken together and considering that in linear heating experiments trapped He is more readily released than spallogenic 3 He (refs. 1-32 and 1-33), it is concluded that only a minor fraction of trapped He initially present has been retained. Thus, precise determination of the components is not possible because of the substantial loss of He in all three cases.

With respect to the different events recorded by the noble gases in the breccia, these findings can be interpreted as follows. The occurrence of solar Ne and Ar indicates that the matrix of the breccia 15015 contains constituents that were exposed to solar irradiation as finely dispersed material before formation of the breccia. The ²¹Ne and ³⁸Ar exposure ages of the matrix and the clast are 280 ± 25 m.y. The deficit of spallogenic ³He indicates a recent degassing event. Compared to solar He, Ne, and Ar in lunar fines, the breccia contains not only concentrations lowered to a variable degree but also severely altered

TABLE 1-VII. - NOBLE GASES RELEASED FROM 15015, 15, TOGETHER WITH CALCULATED EXPOSURE AGES

[All units in 10⁻⁸ cm³ STP/g unless otherwise indicated]

Sample	3 _{Re}	4 _{He}	20 _{Ne}	21 _{Ne}	22 _{Ne.}	36 _{Ar}	38 _{Ar}	40 _{Ar}			
	1	1	Top (15015,1	1 5,20); 3.06 ¤	1 VB	I	1	1			
Total	32.8 ± 1.8	21 400 ± 1000	9620 ± 450	63 ± 3	820 ± 40	11 900 ± 600	2240 ± 110	23 00 ± 1000			
Spallogenic Trapped	≤32.8		37 ± 7 9580 ± 450	39 ± 6 24 ± 7	42 ± 7 780 ± 43	 11 890 ± 600	 ·2220 ± 110				
Exposure ages, m.y.	≤33			250 ± 40							
Middle (15015,15,14); 3.04 mg											
Total	68 ± 4	167 000 ± 9000	24 900 ± 1300	104 ± 7	2030 ± 120	Lost	••				
Spallogenic Trapped	<u><</u> 68.		40 ± 20 24 900 ± 1300	42 ± 19 62 ± 20	46 ± 20 1990 ± 122						
Exposure ages, m.y.	≤ 68	T	••	270 ± 120	 '		•••				
	-		Bottom (15015,15	,4); 3.35 mg				-			
Total	100 ± 4	274 000 ± 10 000	45 000 ± 1600	160 ± 8	3620 ± 150	24 500 ± 900	4640. ± 200	23 900 ± 1000			
Spallogenic Trapped	<100 		47 ± 11 45 000 ± 1600	59 ± 10 111 ± 13	54 ± 11 ^s 3570 ± 150	: 24 500	' 4570 ± 250				
Exposure ages, c.y.	100		••	310 ± 70	r-	••					
			Clast (15015,15,	27); 16.57 mg		-					
Total	23.0 ± 0.3	29 [.] 300 ± 300	251 ± 3	41.2 ± 0.6	60.9 ± 0.9	169 ± 2	56.6 ± 0.9	5600 ± 250			
Spallogenic Trapped Radiogenic	<u><</u> 23 	 	38.7 ± 2.7 212 ± 5 	40.7 ± 2.0 0.5 ± 2.0	44.8 ± 3.1 16.1 ± 3.2	20.0 ± 1.7 149 ± 2.6	30.8 ± 2.2 27,8 ± 2.5	 - 5400			
Exposure ages m.y. ^a	<u><</u> 23			260 ± 12			^b 203 ± 56 290 ± 20				

^aProduction rates from Bogard et al. (ref. 1-30), but P(38) = 1.4×10^{-8} cm³ STP (g Ca)⁻¹ m.y.¹.

^bProduction rated according to Weber (ref. 1-31).

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28

LUNAR SAMPLE STUDIES

abundance patterns. The bottom sample shows the least depletion and the clast, the largest.

It is interesting that the degree of depletion is paralleled by an alteration of the clement abundances. This change in the element abundances shows that the varying concentrations are not solely caused by differences in matrix grain size. It is tempting to invoke thermal differences as the reason for the solar gas pattern observed. The loss of spallogenic ³He seems to be larger in samples closer to the top of the rock. A concentration gradient is also observed for trapped noble gases. The loss of noble gases may have occurred during the heating caused by the coating of the rock with glass. However, comparison of the noble gas abundance patterns in 15015 to those in lunar fines shows larger variations than those observed during linear heating experiments. This may indicate that shock events played an essential role in. the alteration of the initial abundance patterns.

Carbon, Nitrogen, and Sulfur

Three small samples of the breccia from the top, middle, and bottom of the rock were analyzed for C, S, and N (table 1-VIII). The value for C is in the range of other measurements of C content for Apollo 15 samples (ref. 1-34). Little variation was found, and the average value for three samples (table 1-VIII) is 121 parts per million (ppm). This C content is only 29 ppm lower than the measured value of four fines from the same station (station LM, ref. 1-34). The average S content of the three samples is 630 ppm and is not significantly different from the S content of the fines at the same station (ref. 1-35). The average N content of the three samples is 54 ppm; approximately half the N content measured in the fines (refs. 1-35 and 1-36). The C/N ratio in the breccia is 2.24, whereas in the fines it is 1.41.

All three elements show isotopic depletion of the heavy isotopes relative to the measured values for fines samples 15012 and 15013 from the same station. Fines average $\delta C^{13} = +11$ per mill; breccia = -4.3 per mill; δS^{34} fines = +8.1 per mill; breccia.= -+7.1 per mill; δN^{15} fines = +35.6 per mill; breccia = +0.9 per mill. The greatest difference between isotopic values in the fines and breccias is for N, followed by C, with the least change in S.

Petrographic and microscopic studies (previously discussed) indicate that coincident with, or subsequent to, formation of the breccia, thermal annealing may have occurred. The conclusion that heating occurred is also supported by the very low content of ⁴He. The temperature of the sample heating can be estimated by comparing pyrolysis profiles obtained in the experiments on fines sample 15012 (ref. 1-35). It is evident from this study that, at 1073 K (800° C), >90 percent He, 45 percent N₂, and 20 percent C are lost from the sample. At this temperature, 3 is negligibly affected. Furthermore, C preferentially enriched in C¹³ and N enriched in N¹⁵ are released at temperatures of heating from 873 to 1073 K (600° to 800° C). The relatively greater losses of N_2 from the top of the tock (samples closest to 15015,15,17) suggest that this section may have undergone heating to a slightly higher temperature or remained heated for a longer period.

TABLE 1-VIII.- CARBON, SULFUR, AND NITROGEN IN BRECCIA 15015, OBTAINED EY COMBUSTION IN PARTIAL OXYGEN ATMOSPHERE

Sample no.	Location in consortium slab	Weight, B	C, ppm	6C ¹³ PDB (a)	S, ppm	_б с ³⁴ С.р. (б)	N, PPm	6N ¹⁵ atr	He, ppm
15015,15,17 15015,15,7 15015,15,3	Top Middle Bottom	0.5568 .6393 .4607	131 121 110	^c -2.8 -4.5 -5.5	673 586 628	7.4 7.6 6.3	42 61 59	•1.5 3.2 (e)	d _{, t} <1 -1
Average			121	-4.3	630	7.1	54	.9	

^aPDB - Pee Dee belemnite.

^bC.D. = Canyon Diablo.

^CCO₂ repurified.

^dBelow detection limit.

^e Impure.

Methane and Carbide

Interior samples from the top, middle, and bottom of the slab were examined by dissolution in deuterium chloride (DCl) (38 percent in deuterium oxide); the CH4 and CD4 released were analyzed quantitatively by gas chromatography (table 1-IX). The amount of carbide (hydrolyzable C) indicated by the CD_{A} is essentially the same in each sample; however, the CH₄ (methane trapped gas) concentration is much less in samples from the highly vesicular top portion and from the center of the rock. This observation is paralleled by the light rare-gas concentrations and total N measurements (discussed earlier). The ratio of volatile to involatile C (CD_{4}/CH_{4} ratio; bottom 7.3, middle 26.0, and top ∞) suggests that a preferential loss of the volatile species CH₄ from the uppermost sections of 15015 occurred and that presumably this rock has experienced a thermal gradient.

These results are consistent with the following history for 15015. The rock is a metamorphic breccia; urlike the Apollo 14 metamorphic breccias, which were formed from a soil containing only a small portion of mature regolith (ref. 1-37), it was compacted from very mature fines.

The material constituting the top of 15015 has experienced a process more violent (higher temperatures or longer time) than during the formation of highly vesicular glassy agglutinates. Aliquots of such particles selected by microscopic appearance or by density/magnetic properties from the lunar fines show considerable fractionation of volatile CH_A relative to carbide; CD_4/CH_4 ratios up to approximately 21 have been observed (refs. 1-38 and 1-39). The highly vesicular appearance of the top surface of 15015 is strongly indicative of a thermal mechanism rather than a shock process for the removal of volatile species. The materials comprising the interior and bottom of 15015 have progressively less violent histories. Even so, the middle underwent a process similar to agglutination and the bottom, an event like that usually accompanying the formation of group 0 to 1 regolith breccias or soil microbreccias. Unmetanorphosed breccias exhibit low CD_4/CH_4 ratios (approximately 4.0) approximately equal to the ratios observed for bulk soil and constant with volatile- CH_4 retention (refs. 1-38 and 1-39).

Formation of 15015 from local soil (15041; CD4: 23.5 μ g/g; CH₄: 4.6 μ g/g) by lithification at a temperature of approximately 1073 K (800° C) would result in the observed reduction in the carbide content (ref. 140). Indeed, the material constituting the matrix of the breccia has a bulk chemistry that is identical to that of the local soil (15041) (ref. 1-11); thus, 15015 niay have been formed by an impact occurring at the Apollo 15 site. The similarity in the absolute amounts of carbide for the three sections of 15015 suggests that volatile fractionation resulted from the top section of the rock being at an elevated temperature for a longer time (i.e., the bottom of the rock had a more rapid rate of cooling) rather than from an increased temperature. The circumstances leading to the observed temperature gradient within 15015 arc presently not understood.

TABLE 1-IX QUA	NTITIES OF M	ETHANE AND	DEUTERIUM	CARBIDE	RELEASED	BY
	DISSOLUTION	IN DÉUTERI	IUM CHLORII	DE		

Sample no.	Location in consortium slab	Weight, g	CD ₄ , µg/g	CH ₄ , µg/g	CD4/CH4
	_				
15015,15,21	Тор	0.0728	14.6	<0.01	69
15015,15,21	Тор	.0573	15.3	<.01	80
15015,15,6	Middle	.0098	16.6	.64	26
15015,15,2A	Bottom	.0082	15.0	2.0	7.3

THE HISTORY OF LUNAR BRECCIA 15015

X-Ray Photoelectron Spectroscopic Studies

Three types of sample surface from two chips of rock, 1:015,15,8 and 15015,15,29, were examined. These were an exterior glass face from the bottom of the rock and interior chipped and sawn surfaces.

Intense signals due to oxygen (O) and C were obtained in the XPS analysis of the exterior surface. Other peaks identified included Ca, silicon, (Si). Al, Fe, and Mg. The presence of all elements was confirmed by detailed high-resolution spectra from the appropriate kinetic energy regions of the spectrum. Etching at 1333.22^{-5} N/m² (10^{-5} torr) for 20 minutes using Ar-ion bombardment (removal of very approximately 2 nm (20 Å of material) reduced the C signal to a level just above the background noise and constant with a slow buildup of contamination from residual gases in the spectrometer. The major C peak in the unetched sample was presumably due to surface contamination.

The analysis of a chipped surface (greater in area than the collection area of the spectrometer to ensure maximum sensitivity) again revealed a spectrum dominated by C. This C signal was removed by 30 minutes etching with Ar ions. An examination of cut faces yielded similar results, except in one case when the C/O ratio suggested that contamination from the diamond saw was occurring.

No evidence for the presence of the carbidic C giving rise to CD_4 on DC1 dissolution (discussed earlier) was found, although XPS evidence for this species has been reported for fines and a breccia from Apollo 16 (ref. 1-41). It was also impossible to determine the valence state of $\frac{1}{2}$ from either its peak profile or position, even in cases when the etching procedure enhanced the signal due to Fe. Detailed high-resolution study of the appropriate region of the spectrum also failed to reveal the presence of the N that was observed by pyrolysis (discussed previously).

In conclusion, it appears that XPS is not sufficiently sensitive, at present, to detect the low concentrations of C, free Fe, and N that occur in lunar fines and breecias.

CONCLUSIONS

The consortium investigation allows the following tentative history for 15015 to be drawn.

Breccia 15015 is petrologically assigned to Warner's group 2. It was formed from a well-mixed soil; the many fragment types recognized suggest an origin from a shallow depth of regolith. The bulk chemistry of the matrix is similar to that of the local soil, and presumably 15015 was derived at the Apollo 15 site rather than being deposited there from-elsewhere. The soil here could itself still have a large component of disintegrated "ray" material even though the breccia was generated at the Apollo 15 site.

The varying exposure ages of both clasts and matrix indicate a complex history before lithification. The presence of large amounts of noble gases, carbon compounds, and nitrogen, together with etchable tracks and microcrystallites, suggests a considerable exposure of the constituent grains to solar wind.

The ⁴⁰Ar retention ages and ³⁸Ar exposure ages considered together indicate that 15015 was not consolidated earlier than 2.7 ± 0.2 b.y. However, the 0.8to 1 2-b.y. age of the frothy vesicular glass estimated from a ⁴⁰Ar/³⁹Ar "isochron" very tentatively suggests that lithification occurred considerably later. During consolidation, the maximum temperature reached was approximately 1073 K (800°C) based on carbon chemistry and noble gas measurements. Both suggest, however, that 15015 exhibits a unique thermal gradient and that the top of the rock was maintained at a higher temperature or was heated considerably longer than the bottom. Shock events may also have played a part in the preferential fractionation of volatile components from the upper regions of the breccia.

Helium-3 and the track density in the coating glass indicate that after consolidation 15015 was buried at a depth greater than 2 m until it was ejected onto the lunar surface possibly as recently as 30 m.y. ago.

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31

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2. Composition of Lunar Basalts 10069, 10071, and 12008

E. Jarosewich^a and Brian Mason^a....

Chemical analyses, Cross-Iddings-Pirsson-Washington norms, and mineralogical and petrographical data are presented for lunar basalts 10069 and 10071 (from the Apollo 11 mission) and 12008 (from the Apollo 12 mission).

As part of the program for filling gaps in the analytical base for some of the early Apollo missions, samples of basaltic rocks 10069, 10071, and 12008, collected during the Apollo 11 and 12 missions, were allocated for analysis. The results are reported in table 2-I with their Cross-Iddings-Pirsson-Washington (CIPW) norms.

DISCUSSION

Approximately 0.5 g of sample was taken for each chemical analysis. Silica, magnesia (MgO), lime (CaO), total iron (Fe) as ferrous oxide (FeO), titania (TiO₂), and manganese oxide (MnO) were determined by using the general procedure for silicate analysis (ref. 2-1). Aluminum (Al) was determined gravimetrically by precipitation with 8-hydroxyquinoline on an aliquot from the R_2O_3 solution after removal of interfering ions by means of the mercury electrode. Sodium (Na) and potassium (K) were determined by flame photometry; and phosphorus (P) and chromium (Cr) were determined colorimetrically with molybdenum blue (ref. 2-2) and diphenyl carbazide

TABLE 2-I.- CHEMICAL ANALYSES AND CIPW NORMS OF LUNAR BASALTS⁸

Compound	Sample no.					
or mineral (b)	10069	10071	12008			
Сопроч	d abundance	e, wt.X				
S102	40.41	40.47	42.58			
T10,	11.97	12.31	4.82			
A1203	8.22	8.08	8.49			
Cr203	. 36	. 33	.53			
Fe0 Mn0 Mg0 Ca0 Na ₂ 0 K ₂ 0 P ₂ 0 ₅ Total	19.76 .27 7.75 10.50 .55 .31 .21	19.79 .26 7.56 10.63 .54 .32 .18	22.40 .28 11.99 9.10 .24 .14 .05			
CI		.2	100102			
	w norm, w					
Quartz Orthoclase Albite Anorthite Diopside Hypersthene Olivine Chromite Ilmenite Apatite Fem ^C	2.08 1.84 4.66 18.94 26.09 22.61 .53 22.70 .50 .40	2.28 1.89 4.56 18.63 27.08 21.69 	.83 2.04 21.65 19.26 27.73 19.02 .78 4.13 .10 .46			

^aE. Jarosewich, analyst.

 b SiO₂ = silica, Al₂O₃ = alumina,

 Cr_2O_3 = chromic oxide, Na_2O = soda, K_2O = potash, and P_2O_5 = phosphorus pentoxide.

^CFem = FeO/FeO+MgO mole ratio in normative silicates.

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^aSmlthsonian Institution.

(ref. 2-3), respectively. Summations are somewhat greater than 100.00: this feature has been noted by other analysts of lunar rocks. Some of this excess can be ascribed to the presence of metallic Fe, whereas all Fe is reported as FeO. It is possible that titanium (Ti) and Cr may be present in lower valence states than reported. It was noted that ignition of the samples in air results in a weight gain, which is evidently the result of combination with oxygen (O). To economize on the consumption of material, sulfur (S) was not determined. Similar Apollo 11 rocks average 0.2 percent S, and 12051, a rock similar to 12008, contains 0.09 percent S (ref. 2-4).

ROCK_SAMPLES 10069, 10071

Schmitt et al. (ref. 2-5) describe 10069 as a 120-g specimen of very fine grained, vuggy to vesicular, glass-bearing granular basalt aggregate (type A), and 10071 as a 195-g specimen of fine-grained, vesicular to vuggy, glass-bearing plumose basalt aggregate (type AB). Basically, these are hand-specimen descriptions. James and Jackson (ref. 2-6) have provided more detailed descriptions based on thin-section examinations. They classify both rocks as intersertal basalts - 10069 as a very fine grained hornfels, and 10071 as unmetamorphosed. Their modal analysis of 10069 by volume percentage is pyroxene, 54.3; plagioclase, 22.4; ilmenite, 14.9; troilite, 0.9; cristobalite, 4.1; myrmekite and residual glass, 3.2; and apatite, 0.2. For 10071, the modal analysis is olivine, 1.7; pyroxene, 51.5; plagioclase, 20.6; ilmenite, 16.8; armalcolite, trace; troilite, 0.1; cristobalite, 2.8; nivrmekite and residual glass, 6.2; apatite, trace; and others, 0.1. The chemical analyses (table 2-1) show that the two rocks are essentially identical in composition. The potash (K_2O) content shows that the rocks belong to the high K, rubidium (Rb), zirconium (Zr), barium (Ba), cerium (Ce), thorium (Th), uranium (U) geochemical group of the Apollo 11 basalts (ref. 2-7, p. 120). A noteworthy feature is the presence of quartz in the norms and cristobalite in the modes despite the low silicon dioxide (SiO_2) content This feature is related to the high TiO_2 content. which combines with much of the FeO as ilmenite.

A considerable amount of chemical data has been published concerning these two rocks. Major element data for 10069 arc given by Ehmann and Morgan (ref. 2-8), Goles et al. (ref. 2-9), and Teta et al. (2-10). Minor and trace element data are provided by Goles et al. (ref. 2-9), Tera et al. (ref. 2-10), Annell and Helz (ref. 2-11), Murthy et al. (ref. 2-12), and Lovering and Butterfield (ref. 2-13). 1

Major element data for 10071 are given by Ehmann and Morgan (ref. 2-8) and Goles et al. (ref. 2-9); minor and trace element data. by Goles et al. (ref. 2-9), Annell and Helz (ref. 2-11), Gast et al. (ref. 2.14), and Tatsumoto (ref. 2.15). These data are ccmpiled in tables 2.11 and 2.111. For sample 100(59, marked discrepancies exist between the data of the authors of this paper and those of Ehmann and Morgan, and Goles et al. for O, magnesium (Mg), Al, and Fe. In all instances, the authors' measurements are greater than those obtained by instrumental neutron activation analysis by the other investigators. The authors believe their data are to be preferred. because they alone provide a summation close to 100 percent. The large discrepancy between the oxygen data seems certainly to be due to an error in the instrumental analysis, because Ehmann and Morgan record 40.3 percent O in 10071, a rock essentially identical to 10069. Agreement between the authors' measurements and those of other investigators for 10071 is good to excellent.

ROCK SAMPLE 12008

Rock 12008 is a 58-g specimen collected along the traverse route between the lunar module and Middle Crescent crater during the first extravehicular activity on the Apollo 12 mission. Warner (ref. 2-16) classifies rock 12008 along with 12009 and 12015 as type I of the group I lunar basalts. These are porphyritic basalts with skeletal and euhedral olivine phenocrysts in a dark, glassy matrix and have been described as olivine vitrophyres. A polished thin section (fig. 2-1) provided with the analyzed sample shows equant and prismatic olivine phenocrysts (0.1 to 0.2 mm in cross section, up to 0.6 mm long) in a black, opaque groundmass that contains dendritic skeletal olivine crystals (up to 1 nim long, but only 0.03 mm across) and some red-brown titanaugite with a feathery texture suggesting formation by devitrification. The rock contains 21 volume percent olivine phenocrysts, determined by point counting. The compositions of the interiors of the olivine phenocrysts range from 28 to

COMPOSITION OF LUNAR BASALTS 10069, 10071, AND 12008

	Abundance as decormined by -							
Elemont	Jaronewich and Mason	Ehmann and Morgan (rof. 2-8)	Goles st al. (ref. 2+9)	Tera et al. (ref. 2-10)	Annell and Helz (ref. 2-11)	Murthy ot al. (ref. 2-12)	Lovering and Autterfield (ref, 2–13)	
Lithium, ppm	••	••	••	17.2	18	••	••	
Boryllium, ppm · · · ·	••			••	3.3	••	••	
Boron, ppm	••	••	••	••	<10	••	••	
Oxygen, percent	40.83	37.6	••	••	••	••.	••	
Sodium, ppm	4100	••	3650	3400	••	••	••	
Magnesium, percent	4.67	••	3.7	••		••	••	
Aluminum, percent	4.35	3.7	3.8	••	••	••	••	
Silicon, percent	18.89	18.3	18.3	••	••	••	••	
Phosphorus, ppm	900	••	••	••	<2000	••	••	
Potassium, ppm	2600	••		2438		2295	••	
Calcium, percent · · · ·	/.50		7.1	7.20	••	••	••	
Titanium percent	7 10		7 2.4		94			
Vanadium upm			,.2 97					
	2500		2130		2760			
Manganese, ppm	2100		1600		2390	••	••	
Iron, percent	15.36		14.1			••	••	
Cobalt, ppm			26.0		30			
Nickel, ppm			••	••	6.7	••		
Copper, ppm		••	12		8.7		••	
Zinc, ppm				••	<4	••	••	
Gallium, ppm		••	••	••	4.9	••	••	
Germanium, ppm				••	<1	••	••	
Arsenic, ppm				••	< 4		••	
Rubidium, ppm		-•		5.70	5.5	5.60	••	
Strontium, ppm				155.6	1 30	165	••	
Yttrium, ppm				••	164	••	••	
Zirconium, ppm		••	520		566			
N10D1um, ppm	••		••	••	20	••	••	
Molybdenum, ppa				••	<2			
Silver, ppm					<.2 .2			
Indium nom					~ 1			
					<10			
Antimony, ppm					<100			
Tellurium, ppm					< 300			
Cesium, ppm				.163	<1	••		
Barium, ppm			250	288	420	277	••	
Lanthanum, ppm			23.7	••	27	••	••	
Cerium, ppn		••	65		<100	••	••	
Neodymium, ppm					<100		••	
Samailum, ppm			18.0	••	••	••		
Europium,ppm			2.04		••	•-		
Terbium, ppm			4.8	••		••	••	
Kolmium, ppm			6.9		••	••	••	
ttteroium, ppm · · · · ·	••		20.8	••	••	••		
			2.0/	••		••		
			17.0		<20			
Tungeten pom			2.7		< 200			
rungseen, ppm i					~200			
Rhenium, ppm				••	< 30		⁶ 1.18	
Osmilum, ppb			••			••	a. 897	
Platinum, ppm		••	••		<1	••	• •	
Gold, ppm	••			••	<.2	••	**	
Mercury, ppm	••	••		••	- 6		••	
Thallium, ppm	••				- 1	••		
Lead, ppm	••		••	••	-1	••	••	
Bismuth, ppm		••		••	<1		••	
Thorium, ppm		••		••	< 10 0			
Uranium, ppm	••	••	. 78	••	- 500	••	••	

TABLE 2-II- ELEMENTAL ABUNDANCES IN LUNAR BASALT 10069

^aln ppb.

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CONTRACTOR OF THE

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LUNAR SAMPLE STUDIES

Element Jarowski Benann and mand Golu (ref. 2-8) Annell and (ref. 2-1) Gast (ref. 2-1) Gast (ref. 2-14) Lithium, ppn		Abundance as determined by -							
Lithium, ppm. 17 Boron, upm, ppm. 10 Sodium, ppm. 10 Sodium, ppm. Sodium, ppm. Silcon, precent 4.26 Phosphorus, ppm 2700 Calciam, precent 7.60 Calciam, ppm. 7.60 Calciam, ppm. 7.60	Element	Jaronewich and Manon	Elimann and Morgan (ref. 2-8)	Golen et al. (ref. 2-9,	Annell and Helg (ref. 2–11)	Gaar et al. (rof, 2-14)	Tatououto (ref. 2+15)		
Bervilling, pp	Lithium, ppm	C		•/*	17	••	••		
Boron, ppn	Beryllium, ppm			••	3.0	••	••		
Orwein, porcent 40,07 40,03	Boron, ppm	••	••	••	• 10	••	••		
Society pp Society pp <thsociety pp<="" th=""> Society pp Societyp</thsociety>	Oxygen, percent	40.70	40.3	••	••	••	••		
Angeneration process :	Munautum paravat	4000		3040		39803			
Silicon, purcent 19.2 19.1 19.1	Aluminum purcent	4.30	4.2	4.4		••	••		
Phosphorus, ppm 2000	Silicon, percent	18.92	19.1	19.1	••	••	••		
Potamium, ppn 2700 2770 Scandium, ppn 7.2 2770 Scandium, ppn 7.3 97 Scandium, ppn 92 78 Vanadium, ppn 2000 1650 2230 Maganese, ppn 2000 16.9 Cobalt, ppn 7.0 Cobalt, ppn 11 14 Cobalt, ppn <td< td=""><td>Phosphorus, ppm</td><td>800</td><td>••</td><td></td><td>· 2000</td><td>••</td><td></td></td<>	Phosphorus, ppm	800	••		· 2000	••			
Calcium, percent 7.60 7.2 7.2 97 Titanium, percent 7.38 <td< td=""><td>Potassium, ppm</td><td>2700</td><td></td><td></td><td>••</td><td>2770</td><td></td></td<>	Potassium, ppm	2700			••	2770			
Scandium, ppm 73.2 97 Vanadium, ppm 92 78 Vanadium, ppm 2000 1650 2230 Manganese, ppm 2000 1650 2230 Cobalt, ppm 7.0 Cobalt, ppm 7.0 Cobalt, ppm 7.0 Cobalt, ppm 7.0 Cobalt, ppm 4.8 Germanium, ppm 4.8 Strontium, ppm 4.8 Strontium, ppm 162 Zirconus, ppm 162 Cabatum, ppm	Calcium, percent	7.60	••	7.2	••		••		
Titanum percent 7.38 7.00 </td <td>Scandium, ppm</td> <td></td> <td>••</td> <td>73.2</td> <td>97</td> <td>••</td> <td></td>	Scandium, ppm		••	73.2	97	••			
Vanadium, ppm 92 78 Manganese, ppm 2000 1650 2230 Iron, percent 15,38 16,49 Cobalt, ppm 7,0 Copper, ppm 7,0 Callum, ppm 4 Gallum, ppm 4.8	Titanium, percent	7.38		7.0	••	••			
Chromitum, ppm 2000 - 2000 - 1650 2210	Vanadium, ppm		••	92	78	••	••		
Manganese, ppm 2000 1030 2230 1 1 Cobalt, ppm -	Chromium, ppm	2 300		21/0	3060	••	••		
100, percent 17.7 33 Nickel, ppn 7.0 2inc, ppn 11 14 2inc, ppn 4 Copper, ppn 4 Cermanium, ppn 4 Rubidium, ppn 5.2 5.93 Rubidium, ppn 140 160.9 Yttrium, ppn 140 160.9 Yttrium, ppn 140 160.9 Yttrium, ppn 140 160.9 Yttrium, ppn 162 Silver, ppn 162 Indium, ppn 1 Indium, ppn 100 Indium, ppn <td< td=""><td>Manganese, ppm · · · ·</td><td>15 10</td><td></td><td>1650</td><td>2230</td><td></td><td></td></td<>	Manganese, ppm · · · ·	15 10		1650	2230				
Matket, ppm	Cobalt ppm		••	27.1	33	••	••		
Gopper, ppm	Nickel, ppm				7.0	••	••		
Zinc, ppn	Copper, ppm			11	14	••			
Gallium, ppm 4.8 Germanium, ppm Rubidium, ppm Rubidium, ppm 140 160.9 Strontium, ppm 140 160.9 Vttrium, ppm 140 160.9 Vttrium, ppm 140 Nobidenum, ppm 24 Silver, ppm	21nc, ppm				.4				
Germanium, ppm	Gallium, ppm	••			4.8	••			
Arrenic, ppm	Germanium, ppm					••			
Rubidum, ppm	Arsenic, ppm				•4	••			
Strontum, ppm	Rubidium, ppm		••	••	5.2	5.93			
11:1:1:1:1:1:1:1:1:1:1:1:1:1:1:1:1:1:1	Strontlum, ppm				140	160.9			
1100 100 100 100 100 Molbidenum, ppm Silver, ppm Cadmium, ppm Cadmium, ppm Tin, ppm 10 Talium, ppm 100 Cesium, ppm 300 Cesium, ppm 300 Cerium, ppm 300 Cerium, ppm 25.8 27 28.8 Cerium, ppm 20.0 22.7 Europlum, ppm 20.0 22.7 Europlum, ppm 20.0 22.7 <td>Zirconium ppm</td> <td>••</td> <td>••</td> <td>210</td> <td>102 644</td> <td>••</td> <td></td>	Zirconium ppm	••	••	210	102 644	••			
Molybderum, ppm	Nichium, ppm				24				
Silver, ppm	Molybdenum, ppm				<2				
Cadmium, ppm	S11ver, ppm	••			·.2	••			
Indium, ppm	Cadmium, ppm				۰8	••			
Tin, ppm	Indium, ppm					••	••		
Antimory, ppm	Tin, ppm	••		••	• 10	••	••		
1 1 100 100 100 Barium, ppm 1 100 327 100 Barium, ppm 1 100 327 100 Lanthanum, ppm 1 100 83.5 100 Neodynium, ppm 1 100 64.5 100 Samarium, ppm 1 1 100 64.5 100 Samarium, ppm 1 1 100 64.5 100 Samarium, ppm 1 1 1 1 1 1 Gadolinium, ppm 1 <td>Antimony, ppm</td> <td></td> <td></td> <td></td> <td>100</td> <td>••</td> <td></td>	Antimony, ppm				100	••			
Certain, ppm	Castum ppm	••		••	. 300	170			
Latthalum, ppm 25.8 27 28.8 Cerlum, ppm 84 -100 83.5 Neodymium, ppm 20.0 22.7 Europium, ppm 22.7 22.7 Europium, ppm 22.7 22.7 Gadolinium, ppm 22.7 29.3 Terbium, ppm 29.3 Dvsprosium, ppm Dvsprosium, ppm	Barlum, ppm			450	470	327			
Cerium, ppm 84 -100 83.5 Neodymium, ppm 20.0 22.7 Samarium, ppm 20.0 22.7 Gadolinium, ppm 212 23.42 Gadolinium, ppm 29.3 29.3 Terbium, ppm 29.3 Dvsprosium, ppm 29.3 Erbium, ppm 9.2 Erbium, ppm 9.2 Erbium, ppm 20.8 20.5 Hafnium, ppm 2.00 100 Tungsten, ppm	Lanthanum, ppm			25.8	27	28.8			
Neodymium, ppm	Cerium, ppm			84	• 100	83.5			
Samarium, ppm 20.0 22.7 Europium, ppm 2.12 2.32 Gadolinium, ppm 2.7 2.32 Terbium, ppm 2.32 2.32 Terbium, ppm 2.7 2.32 Terbium, ppm 2.62 2.9,3 Dysprosium, ppm 29,3 Dysprosium, ppm 33.5 <td< td=""><td>Neodymium, ppm</td><td></td><td></td><td></td><td>· 100</td><td>64.5</td><td>••</td></td<>	Neodymium, ppm				· 100	64.5	••		
Europium, ppm 2.12 2.62 Gadolinium, ppm 29.3 Dvsprosium, ppm 29.3 Dvsprosium, ppm 33.5 Dvsprosium, ppm 9.2 Terbium, ppm 9.2 Ytterblum, ppm 20.8 2.67 Lutetium, ppm 19.1 20 Tantalum, ppm 100 Rhenium, ppm 100 Tantalum, ppm 100 Rhenium, ppm Tunksten, ppm Hercurv, ppm -	Samarium, ppm			20.0		22.7	••		
Gadol Inium, ppm 29,3 Terbium, ppm 5.7 Dvsprosium, ppm 5.7 Holmium, ppm 9.2 Holmium, ppm 20.8 20.5 Luctium, ppm 20.8 2.67 Hafnium, ppm 2.0 100 Tantalum, ppm Rhenium, ppm 2.0 100 Rhenium, ppm	Europium, ppm	••		2.12		2.42			
Territin, ppm	Gadolinium, ppm	••	••		••	29.3	••		
Dysproslum, ppm Holmium, ppm Erbium, ppm 20.8 Uterbium, ppm 20.8 2.67 Lutetium, ppm 19.1 -20 Tantalum, ppm 19.1 -20 Tantalum, ppm 2.67 Tantalum, ppm 100 Tunksten, ppm Platinum, ppm Wercurv, ppm Thallium, ppm Issauth, ppm Thallium, ppm Issauth, ppm Issauth, ppm Issauth, ppm <	Terbium, ppm					11 6			
Totaling, ppm	Halmium num	••		9.2					
Ytterblum, ppm 20,8 20,5 Lutetium, ppm 3.08 2.67 Hafnium, ppm 19,1 200 Tantalum, ppm 2.0 100 Tunksten, ppm 2.00 Rhenium, ppm Rhenium, ppm Rhenium, ppm Rhenium, ppm Wercurv, ppm	Erbium, ppm		••	**	••	21.3			
Lutetium, ppm 3.08 2.67 Hafnium, ppm 19.1 -20 Tantalum, ppm 19.1 -20 Tantalum, ppm 2.0 100 Tumsten, ppm 200 Rhenium, ppm 30 Gold, ppm Mercurv, ppm <t< td=""><td>Ytterblum, ppm</td><td></td><td></td><td>20.8</td><td>••</td><td>20.5</td><td>••</td></t<>	Ytterblum, ppm			20.8	••	20.5	••		
Hafnium, ppm 19,1 -20 Tantaium, ppm 2.0 100 Tungsten, ppm Rhenium, ppm Platinum, ppm Gold, ppm Wercurv, ppm Thallium, ppm Bismuth, ppm 1 Thorium, ppm 100 4-4.4, 3/24 Uranium, ppm	Lutetium, ppm		••	3.08		2.67			
Tantalum, ppm 2.0 -100 Tunksten, ppm -200 Rhenium, ppm Platinum, ppm Wercurv, ppm Thallium, ppm Isad, ppm Isad, ppm Isad, ppm 1 Isad, ppm 1 Isad, ppm -1 1 Isad, ppm 1 </td <td>Hafnium, ppm</td> <td></td> <td></td> <td>19.1</td> <td>· 20</td> <td></td> <td></td>	Hafnium, ppm			19.1	· 20				
Tunksten, ppm ***	Tantalum, ppm			2.0	· 100	••			
Rhenium, ppm -10 -10 Platinum, ppm -3 Gold, ppm -3 Mercurv, ppm Mercurv, ppm Mercurv, ppm Signuth, ppm Bismuth, ppm Thorium, ppm 100 Uranium, ppm -69 -500 .8/4, .84%	Tunksten, ppm				*200				
riatinum, ppm	Rhenium, ppm				- 30				
tota, ppm	riatinum, ppm · · · ·								
Thallium, ppm	Moreury nam								
Lead, ppm 1 1.71, 1.67 Bismuth, ppm -1 1 Thorium, ppm -100 8.43, 3.24 Uranium, ppm -69 -500 .873, 365	Thallum, nnm				.1		••		
Bismuth, ppm -1 Thorium, ppm -100 6.43, 3.34 Uranium, ppm -69 500 .8/3, 3.34	lead, ppm				. 1	••	1.71, 1.67		
Thorium, ppn	Bismuth, ppm	••	••		- 1	••			
Uranium, ppm	Thorium, ppm				+ 100		4.44. 4.24		
	l'ranium, ppm			.69	+ 500		1874, 1845		

TABLE 2-III.- ELEMENTAL ABUNDANCES IN LUNAR MASALT 10(7)

COMPOSITION OF LUNAR BASALTS 10069, 10071, AND 12008



Figure 2-1.- Photomicrograph (transmitted light) of a thin section of lunar basalt 12008, showing phenocrysts of olivine (maximum length 0.6 mm) in opaque groundmass.

35 ntol% fayalite, with an average of 32 mol%. In reflected light, the groundmass is seen to contain occasional small euhedral chromite grains and rare minute metal particles.

The analysis of 12008 is comparable with analyses of other Apollo 12 basalts, such as 12022 (ref. 2-17), 12051 (ref. 2-4), and 12063 (ref. 2-18). The TiO_2 content (4.82 percent) is somewhat higher than many of the Apollo 12 basalts, the average for which is approximately 3 percent. The amount of normative olivine is somewhat less than the modal amount, indicating that the groundmass may be slightly oversaturated. In the classification of James and Wright (ref. 2-19), 12008 belongs in group IIA, ilmentte-bearing olivine basalts. It is evidently a quenched sample of this subordinate magma type at the Apollo 12 site.

Anders et al. (ref. 2-20) provide trace element data for 15 elements in 12008.

ACKNOWLEDGMENTS

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3. Composition of Eight Apollo 17 Basalts

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Chemical analyses, Cross-Iddings-Pirsson-Washington norms, and mineralogical and petrographical data are presented for eight mare basalts collected on the Apollo 17 mission. Seven of these have very similar chemical compositions but can be divided into two textural groups: medium to coarse grained (70017, 71055, 74255, 75075) and fine grained (70215, 71569, 74275). The eighth, 75035, has a distinctive chemical and mineralogical composition (lower titania, higher silica, quartz in the-norm, and cristobalite in the mode), and evidently represents a different flow probably underlying the hightitanium basalts at Camelot Crater.

The authors received samples of the following Apollo 17 basalts for analysis:

1. Sample 70017: A 2957-g, coarse-grained, vuggy basalt collected close to the lunar module (LM)

2. Sample 70215: An 81-10-g, dense, fine-grained, nonvesicular basalt collected approximately 50 m northeast of the LM

3 Sample 71055: A 669-g, medium-grained, very vesicular basalt collected at station 1a

4. Sample 71569: A 289-g, fine-grained basalt collected at station 1a (rake fragment)

5. Sample 74255: A 737-g, coarse-grained, vuggy basalt collected at station 4 (south rim of Shorty Crater)

6. Sample 74275: A 1493-g, dense, fine-grained basalt collected at station 4

7. Sample 75035: A 1235-g, medium-grained basalt collected at station 5 (southwest rim of Camelot Crater)

8. Sample 75075: A.1008-g, coarse-grained, vuggy basalt-collected at station 5

Approximately 0.5 g of sample was taken for eachof the chemical analyses listed in table 3-I. Silica (SiO_2) , magnesia (MgO), lime (CaO), total iron (Fe) as ferrous oxide (FeO), titania (TiO₂), and manganese monoxide (MnO) were determined according to the general procedure for silicate analysis (ref. 3-1). Alumina (Al_2O_3) was determined gravimetrically by precipitation with 8-hydroxyquinoline on an aliquot from the R₂O₃ solution after removal of interfering ions by means of the mercury electrode. Soda (Na₂O) and potash (K_2O) were determined by flame photometry. and phosphorus (P) and chromium (Cr) were determined colorimetrically with molybdenum blue (ref. 3-2) and diphenyl carbazide (ref. 3-3), respectively. Summations often are somewhat greater than 100.00; this feature has been noted by other analysts of lunar rocks. Some of this excess can be ascribed to the presence of metallic Fe, whereas all Fe is reported as FeO. It is also possible that titanium (Ti) and Cr may be present in lower valence states than reported. Ignition of the samples in air resulted in a weight gain, evidently the result of combination with oxygen (O). To economize on the consumption of material, sulfur (S) was not determined; analyses reported by the Apollo 17 preliminary examination team (ref. 3-4) show about 0.17 percent S in the Apollo 17 basalts, which is equivalent to approximately 0.5 percent troilite.

Modal analyses were made of most of the rocks by the point-counting technique. In general, the modal analyses were in good agreement with the normative composition, which is to be expected in view of the close correspondence between normative and ob-

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Compound	Sample no.											
or mineral (b)	74255	74275	75075	70017	71569	71055	70215	75035				
	Compound abundance, wt.X											
SID,	37.98	38.21	37.45	37.62	38.63	37.30	37.71	41.29				
T10,	12.55	12.42	13.96	13.90	12.75	13.27	13.48	10.40				
A1,0,	8.94	8.62	8.59	8.85	7.99	9.45	9.08	9.52				
Cr.0,	.56	.55	.51	.54	.42	-40	.40	-23				
FeO	18.64	18.67	19.82	1.9.60	20.05	20.00	20.01	18.78				
MnO	.29	.27	.27	.29	.29	.26	.26	.25				
MgO	10.73	10.63	9.58	9.42	8.99	8.44	8.23	6.50				
CaO	10.09	10.19	9.83	9.81	10.31	10.52	10.71	13.07				
Na ₂ 0	.39	. 37	. 37	.41	.39	.43	.42	.46				
к ₂ ō	.08	.08	.06	.07	.ò7	.06	.05	.07				
P20.5	.04	.03	.04	.04	.04	.10	.07	.05				
Total	100.29	100.04	100.48	100.55	99.93	100.23	100.42	100.62				
				CIPW not	m, wt.%							
Quartz					.16			1.60				
Orthoclase	0.47	.47	. 35	.41	.41	. 35	.28	. 39				
Albite	3.30	3.13	3.13	3.47	3.30	3.64	3.51	3.88				
Anorthite	22.41	21.62	21.60	22.10	19.85	23.68	22.71	23.60				
Diopside	22.19	23.29	21.86	21.37	25.40	22.80	24.52	33.91				
Hyperstheme	20.97	21.98	24.97	25.19	25.89	20.86	22.61	16.94				
Olivine	6 21	5.08	1.21	.72		2.88	.37					
Chromite	.82	. 81	.75	.80	.62	.59	.59	.34				
Ilmenite	23.84	23.59	26.51	26.40	24.22	25.20	25.50	19.80				
Apatite	.09	.07	.09	. 09•	.09	.23	.17	.10				
Fém ^C	.28	.28	. 30	. 30	. 35	.35	. 35	.45				

TABLE 3-I.- CHEMICAL ANALYSES AND CIFW NORMS OF APOLLO 17 BASALTS, IN ORDER OF MgO CONTENT⁴

^a. Jarosewich, analyst.

 ${}^{b}Si0_{2} = silica, Ti0_{2} = titania, Al_{2}O_{3} = alumina, Cr_{2}O_{3} = chromic oxide, FeO = ferrous oxide,$ $MnO = manganese monoxide, MgO = magnesia, CaO = lime, Na_{2}O = soda, K_{2}O = potash, P_{2}O_{5} = phosphorus pentoxide..$

^CFem = FeO/FeO+MgO mole ratio in normative silicates.

served inineralogy. Modal pyroxene is a few percent higher than normative pyroxene, because of the incorporation of minor amounts of Al_2O_3 and TiO_2 ; and, because not all Al_2O_3 is combined as plagioclase, the modal amount of the latter is slightly less than the normative amount. When olivine is present in the norm, even in amounts less than I percent, it is seen in thin sections. It is also present as small phenocrysts in 71569, although the analysis is slightly quartz-normative; interstitial cristobalite (or tridymite) was detected in the groundmass with the microprobe. Cristobalite is a prominent accessory in 75035, amounting to about 5 percent, although only 1.6 percent quartz is in the norm; the additional SiO₂ is evidently provided by substitution of aluminum (Al) for silicon (Si) in pyroxene.

Trace element determinations on aliquots of the analyzed samples are reported in table 3-II. The technique used is described by Thompson and Bankston (ref. 3-5). Trace element variations within these basalts are not great, which is consistent with the limited range in the major element composition. Compared to data on Apollo 15 basalts (ref. 3-6), the Apollo 17 basalts have lower Cr, cobalt (Co), and nickel (Ni) and higher gallium (Ga), strontium (Sr), vanadium (V), and zirconium (Zr). The higher V can probably be correlated with the high Ti content of the Apollo 17 rocks.

Element	Sample úo.							
	74255	74275	75075	70017	71569	71055	70215	75035
				Abundan	ce, ppm			
Barium_	130	100	45	45	80	50	55	150
Lead	10	6			; 18			3
Strontium	275	190	120	140	145	130	130	295
Yttrium	1 35	135			145			155
Zirconium -	125	145	320	235	175	240	240	220
Nickel	19	33	20	20	25	28	11	22
Cobalt	12	13	16	23	5	27	21	8
Zinc	<2	<2	5	4	4	5	6	<2
Copper	9	7	30	17	8	25	16	7
Lithium	9	9	9	10	10	13	10	12
Chromium	>2000	>2000	3100	2950	>2000	2950	3050	1500
Vanadium	240	270	290	260	280	315	320	140
Gallium	13	10	13	9	13	11	11	10
Boron	<2	<2	1	1	<2	2	3	<2

TABLE 3-II. - TRACE ELEMENT ANALYSES OF APOLLO 17 BASALTS^a

^aG. Thompson, analyst.

PETROLOGY

Table 3-I shows that the basalts analyzed are remarkably uniform in composition, with the exception of 75035. Omitting the latter from consideration for the present, the remaining seven basalts can be clearly divided into two distinct textural groups as follows.

1. Samples 7001,7, 71055, 74255, and 75075 are medium- to coarse-grained basalts with ophitic to sub-ophitic textures.

2. Samples 70215, 71569, and 74275 are finegrained basalts with acicular ilmenite crystals and skeletal and/or subequant phenocrysts of olivine in a groundmass of plagioc'ase and pyroxene.

Despite the textural differences, the compositions may be essentially identical (as for 74255 and 742.5), suggesting that the differences reflect different cooling histories. Other investigators have also distinquished these two groups of Apollo 17 basalts; group A corresponds to type 3 of Papike and Bence (ref. 3-7) and mare basalts III of Brown et al. (ref. 3-8), whereas group B corresponds to type 2 of Papike and Bence and mare basalts I of Brown et al. Brown et al. classify 75035 as mare basalt II, and it probably belongs to type I of Papike and Bence. It is therefore convenient to describe the petrology of the samples in terms of these three groupings.

ROCK SAMPLES 70017, 71055, 74255, AND 75075

Samples 70017, 71055, 74255, and 75075 are all mediunt-to coarse-grained basalts (maximum length of plagioclase crystals, 1 to 4 mm) with ophitic to subophitic textures. The principal minerals are calcic and subcalcic augite, plagioclase, and ilmenite, with minor amounts (1 to 5 percent) of olivine; some cristobalite is usually present as late crystallization. Trace constituents are armalcolite (in ilmenite), chromian ulvöspinel, troilite, and metallic Fe. Brown, silica-rich (SiO₂~70 percent) and potassium-rich (K₂O~5 percent) mesostasis is present in small amounts. Plagio clase is very uniform in composition. averaging 82 mol% anorthite (An₈₂); some of the larger crystals are weakly zoned, the maximum range noted being from An₈₅ (core) to An₇₇ (margin). Olivine shows a

moderate range in composition 28 to 38 mol% fayalite (Fa_{28} to Fa_{38}); the olivine in 74255 is more magnesian than that in the other rocks; this correlated with the lower fem¹ value (0.28) for this rock (table 3-I). The pyroxene (fig. 3-1) is largely calcic to subcalcic augite, with minor amounts of pigeonite; the pigeonite occurs sometimes as cores to large augite crystals, and sometimes pigeonite and augite are complexly zoned with each other.

Additional information has been obtained regarding Ti and Al concentrations in the pyroxenes in 70017 and 74275. As can be seen in figure 3-2, the Ti: Al ratio (atomic) straidles the 1:2 line, indicating the presence of a CaTiAl₂O₆ component. The pyroxenes in 74275 are notably richer in both Ti and Al than those in 70017; this suggests later crystallization of ilmenite and plagioclase in 74275 than in 70017, whereby the pyroxene in 74275 crystallized from a magma rich in Ti and Al. A few of the pyroxenes in 70017 fall between the 1:2 and 1:1 line, suggesting that a small amount of Ti3 (as the component $CaTiAlSiO_6$) may be present. On the other hand, the pyroxenes in 74275 plot between the 1:2 and the 1:4 line, indicating the presence of the CaAl₂SiO₆ component.

ROCK SAMPLES 70215, 71569, AND 74275

Samples 70215, 71569, and /4275 are basalts with microphenocrysts of olivine and acicular ilmenite in a fine-grained groundmass of augite and plagioclase. They are indistinguishable in bulk chemistry from the preceding group. The compositions of their constituent minerals are very uniform (fig. 3-3). Olivine ranges from Fa₂₁ to Fa₃₄; individual crystals are sometimes zoned from magnesium-rich cores to more iron-rich margins. The pyroxene is a strongly colored reddish-brown augite of remarkably uniform composition. (Most compositions are within a field 37 to 47 mol% Wollastonite (Wo_{37} to Wo_{47}), 16 to 22 mol% feirosilite (Fs₁₆ to Fs₂₂). Plagioclase occurs as small laths in the groundmass; its composition is also very uniform, An₇₇ to An₈₀. Some of the olivine includes small crystals of chromian ulvöspinel. A little tridymite or cristobalite was detected in the groundmass with the microprobe. Among these three rocks, 70215 is texturally distinctive for the presence of long (up to 2 mm) skeletal clivine crystals.



Figure 3-1.- Composition of olivine and pyroxenes in mare basalts 70017, 71055, and 75075.

¹Fem = Fe0/Fe0+Mg0 mole ratio in normative solicates.

COMPOSITION OF EIGHT_APOLLO 17 BASALTS

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Figure 3-2.- TiO2 and Al2O3 content of clinopyroxenes in mare basalts 70017 and 74275.





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LUNAR SAMPLE STUDIES

ROCK SAMPLE 75035

Rock 75035 is a medium-grained (augite grains up to 0.6 mm across, plagioclase laths up to 1.5 mm long), ophitic, vesicular basalt from the southwest rim of Camelot Crater. near the center of the Taurus-Littrow valley. The primary assemblage is strongly zoned calcic to subcalcic augite (fig. 3-4), plagioclase (unzoned to weakly zoned, ranging in composition from An₈₆ in the centers of large crystals to An₇₇ at their margins), and ilmenite in lathy to blocky crystals. Olivine is absent, and cristobalite is a prominent accessory (~5 percent). The pyroxenes are zoned toward increasing Fe and somewhat diminishing calcium (Ca), going from cores to margins; the zoning does not appear in thin section as discrete bands but as rounded, irregular patches. Many of the pyroxene crystals are rimmed by pyroxferroite. Meyer and Boctor (ref. 3-9) give a detailed account, with analyses, of the opaque minerals in 75035; ilmenite is the major opaque mineral, but ulvöspinel, metallic Fe, troilite, tranquillityite, baddeleyite, and zirconolite are also present in small amounts.

Sample 75035 is unique among the basalts analyzed by the author. Its fem value of 0.45 (table 3-I) is considerably higher than the other basalts, and it has the lowest TiO_2 and MgO and the highest CaO contents. Its mineralogy is notable for the absence of olivine, the presence of appreciable cristobalite, and

the presence of pyroxferroite. All these features suggest a more strongly fractionated magma as the source material.

DISCUSSION

Papike and Bence (ref. 3-7) comment that their type 2 and type 3 basalts, equivalent respectively to the B and A groups of this paper, have essentially identical chemistry and could represent different cooling histories in the same basalt flow. This observation is certainly true for 74255 and 74275, as can be seen from tables 3-I and 3-II. All the major elements and most of the minor elements are identical within the range of probable error. The only exceptions are Ni (1.9 and 33 ppm) and Sr (275 and 190 ppm); the Ni values are probably controlled by the erratic occurrence of minute metal particles, but the cause for the variation in Sr is not obvious. If the Sr variation is dismissed as a chance artifact, then 74255 and 74275 may well be parts of the same flow, 74275 representing a rapidly crystallized marginal phase. These two rocks were collected close together on the rim of Shorty Crater.

The remaining rocks of groups A and B (70017, 70215, 71055, 71569, and 75075) were collected either at, or within approximately 1 km of, the LM, in the central part of the Taurus-Littrow valley. They show some range in major element chemistry, the



Figure 3-4. Composition of pyroxenes and pyroxferrolte in mare basalt 75035.

principal variable being MgO (9.58 to 8.23 percent), resulting in fem values from 0.30 to 0.35. Most of the trace elements are relatively uniform throughout, exceptions being Co (5 to 27 ppm), copper (Cu; 8 to 30 ppm), Ni (11 to 28 ppm), and Zr (175 to 320 ppm). The variations in these trace elements seem random and unrelated to the MgO and fem variation. Cobalt, copper, and nickel are probably concentrated in fare metal particles, and Zr, in an accessory mineral such as baddeleyite; the variation may be more apparent (affected by sampling), rather than a significant effect. All these samples could well be fragments from a single flow or a sequence of thin flows of similar composition.

Rock 75035 is clearly a basalt of distinctive composition. The only comparable rock thus far described from the Apollo 17 mission is 75055, from the same location, the analysis of which (ref. 3-4) is essentially identical. It thus appears that Camelot Crater excavated both high-titanium basalts (75075, 13.96 percent TiO₂) and low-titanium basalts (75035, 10.40 percent TiO₂). Camelot Crater is one of the largest craters in the Taurus-Littrow valley, and it may have penetrated a greater depth of the mare lava flows than the other craters visited. Such an occurrence would suggest that 75035 and 75055 are samples of a deeper, older lava than the basalts of groups A and B.

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4. Troctolitic and Basaltic Clasts from a Fra Mauro Breccia

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A troctolitic clast removed from breccia 14321 is a well-equilibrated subsolidus mineral assemblage of olivine (86 mol% forsterite) and plagioclase (95 mol% anorthite) with accessory whitlockite, chrome spinel, ilmenite, armalcolite, and barian potassium-rich feldspar. This clast is a low-temperature-equilibrated example of the spinel troctolite compositional grouping and contrasts with the high-temperature, quenched mineral assemblages of most rocks in this group. Two basaltic clasts from the same breccia have affinities with rock 14053.

Lunar sample 14321 is a partly annealed but coherent clastic breccia found near the rim of Cone Crater in the Fra Mauro region; it is included in the classification of Chao, Minkin, and Best (ref. 4-1) as a "moderately annealed Fra Mauro breccia" and in the classification of Warner (ref. 4-2) as a "medium grade partly annealed breccia." Grieve et al. (ref. 4-3) have examined the petrology in detail and identify several generations of brecciation culminating in the incorporation of basaltic and other clasts, including anorthosite, in an unrecrystallized light friable matrix.

A troctolitic clast weighing 0.5 g and two basaltic fragments weighing 0.15 g each were removed from the light matrix in consortium sample 14321,88 (W. Compston, consortium leader). The rare-earth element (REE) abundances have been determined by Taylor et al. (ref. 4-4): the basaltic clasts are enriched by factors of 100 to 50 (lanthanum to ytterbium (La/Yb)) relative to chondrites with a moderate negative europium anomaly, and the troctolite is enriched by corresponding factors of 50 to 12 (La/Yb) with a positive europium anomaly. Compston et al. (ref. 4-5) report rubidium to strontium (Rb/Sr) mineral ages of 4.08 ± 0.1 and 4.05 ± 0.08 billion years (b.y.) for the basalts and 3.74 ± 0.17 b.y. for the troctolite.

TECHNIQUES

A 5- by 4-mm polished thin section and a 0.05-g fused glass bead were made from the troctolite fragment. After using the small basaltic clasts for Rb/Sr mineral age determination and for spark source mass spectrography, insufficient material remained for preparation of thin sections; therefore, electron-probe analysis was performed on selected grains from mineral separates.

All electron-probe analyses reported herein were performed using an Applied Research Laboratories EMX microanalyzer and techniques described by Lovering and Ware (ref. 4-6). Measurements also were made using a Technisch Physische Dienst microprobe fitted with a silicon (lithium-activated) detector (ref. 4-7); these additional data confirmed the earlier conventional probe analyses.

TROCTOLITE CLAST 8A

The Cross-Iddings-Pirsson-Washington (CIPW) norm calculated from the electron-probe analysis of the fused glass bead from troctolite clast 8A of sample 14321,88 is given in table 4-I. The CIPW norm is in good agreement with the observed mineralogy in

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LUNAR SAMPLE STUDIES

TABLE 4-1.- COMPARISON OF TROCTOLITE CLAST 8A AND AVERAGE SPINEL TROCTOLITE

Compound or mineral	Troctolite clast 8A	Spinel troctolite							
Compound abundance, wt.%									
Silicon dioxide (SiO ₂)	43.50	43-7							
Titanium dioxide (Ti0 ₂)	. 1.9	.17							
Aluminum oxide (Al ₂ 0 ₃)	23.29	22.7							
Iron oxide (FeO) Manganese oxide (MnO) Magnesium oxide (MgO) Calcium oxide (CaO) Sodium oxide (Na ₂ O) Potassium oxide (K ₂ O) Chromium oxide (Cr ₂ O ₃) Phosphorus pentoxide (P ₂ O ₅) Total	4.56 .06 15.82 12.27 .28 .06 .03 100.06	4.9 .07 14.7 13.1 .39. .05 .11 .04 99.93							
(CIPW norm, wt.%								
Corundum Orthoclase Albite Anorthite Olivine Diopside Hypersthene Ilmenite Chromite	0.5 .4 2.4 60.8 29.7 0 5.9 .4 .04	0 .3 3.4 58.1 31.2 3.6 3.0 .3 .1							

^aSpinel troctolite of Prinz et al. (ref. 4-8).

the thin section, which contains approximately 35 percent olivine (86 mol% forsterite (Fo₈₆), table 4-II) set in plagioclase (95 mol% anorthite (An₉₅), table 4-III). Whitlockite (table 4-IV), which constitutes approximately 1 percent by volume and occurs in groups of 50- μ m cuhedral grains within the plagioclase, is the main accessory phase. Lesser amounts of ilmenite and aluminous chrome spinel are distributed

randomly throughout the clast, and these grains of armalcolite were found associated with ilmenite (fig. 4-1, table 4-V). A few 40- μ m anhedral grains within plagioclase were identified as barian potassium-rich feldspar (K-feldspar) (table 4-III).

The plagioclase, the whitlockite, and the ilmenite are remarkably homogeneous and unzoned, and the compositional scatter in the

TROCTOLITIC AND BASALTIC CLASTS FROM A FRA MAURO BPECCIA

Compound,	Basalt	clasts	Troctolite and basalt fusion crust			
or mineral	4۸	6A	Clast 8A	Glassy basaltic fragment	Contact zone rim ^a	
		Compound	abundance, wi	t.%		
Śi0 ₂	36.05	37.05	40.06	35.29	35.39	
FeO	36.23	29.63	12.89	35.86	37.20	
MnO	.28	. 31	.13	.27	.33	
MgO	27.52	32.73	46.39	28.52	26.54	
CaO	.15	. 14	.03	<.02	.17	
^{Cr} 2 ⁰ 3	.20	.22	<.03	<.03	<.03	
Total	100.43	100.08	99.50	99.94	99.63	
No. grains	1	7	14	1	3	
6442		Molecul	ar proportion	b		
Si	1.001	0.999	1.000	0.985	0.998	
Fe	. 84 1	.668	.269	. 837	.877	
Mn	.007	.007	• .003	.006	.008	
Mg	1.139	1.316	1.727	1.187	1.115	
Ca	.005	.004	.001	.0	.005	
Cr	.004	.005	.0	.0	.0	
		Mineral	abundance, mo	01%		
Total Fo	57.5	66.3	86.5	58.6	56.0	
Max. Fo		68.0	87.1			
Min. Fo		63.9	85.1	••		

TABLE 4-II. - OLIVINE ANALYSES

^aAnalysis at rim to glass.

^bOn the basis of four oxygen ions.

spinel analyses is considerably less than that found in aluminous lunar chromites from other rocks. A small but significant compositional scatter of olivine is evident, but zoning was not observed within individual grains excepting those, described later, at the edge of the clast.

The presence of magnesian armalcolite is unusual: this mineral is usually formed as a liquidus phase in titanium-rich magmas, and any associated chrome spinel has a very high titanium content reflecting the concentration in the liquid. For example, the primary spinel in the armalcolite-bearing rock 70035 contains as much as 23 percent titanium dioxide (TiO₂). The olivine (86 mol% magnesium (Mg₈₆)), armalcolite (Mg_{61.5}), spinel (Mg₅₃), and ilmenite (Mg_{37.5}) association in the troctolite is a subsolidus-equilibrated assemblage that is believed by the authors to be unique among the anorthosite and troctolite samples investigated.

LUNAR SAMPLE STUDIES

Compound.	Basalt	clasts	Troctolite	Glassy						
element, or mineral	4A 6A		Plagioclase	K-feldspar	basaltic fragment					
Compound abundance, wt.%										
\$10 ₂	47.24	46.87	44.91	63.01	47.87					
A1203	33.14	33.55	34.94 _	19.98	32.40					
Fe0	.49	.45	.07	<.03	.53					
MgO	.18	. 19	.04	<.03	. 49					
Ca0	17.54	17.71	19.20	.45	17.02					
Sr0_	<.04	<.04	<.04	.04	<.04					
Ba0 ^a	<.04	<.04	<.04	3.00	<.04					
Na ₂ 0	1.14	1.13	.45	.45	1.63					
к ₂ 0	.15	. 13	.08	13.15	.29					
Total	99 . 88	100.03	99.69	100.08	100.23					
No. grains	23	10	15	3	1					
		Molecular	proportion ^b							
Si	8.701	8.26?	8.318	11.733	8.794					
A1	7.194	7.278	7.627	4.385	7.015					
Fe	.076	.969	.011	.0	.081					
Mg	.049	.052	.011	.0	.134					
Ca	3.461	3.492	3.810	.090	3.350					
Sr	.Ü	.0	.0	.004	.Ò					
Ba	.0	.0	.0	.219	•0					
Na	. 407	.403	. 162	.163	.581					
К	.035	.031	.า19	3.124	.068					
Total	19.923	19.592	19.958	19.718	20.023					
		Mineral ab	oundance, mol%							
Total An	88.7	88.9	95.5	2.5	83.8					
Total Or ^C	.9	.8	.5	86.9	1.7					
Max. An	91.8	93.4	95.7	••	••					
Min. An	80.0	85.0	95.0	••	••					

TABLE 4-III. - FELDSPAR ANALYSES

^aBaO = barium oxide.

 $^{\rm b}{\rm On}$ the basis of 32 oxygen ions.

^COr = orthoclase.

Compound or clement	Abundance, we.2							
Primary dotections								
P ₂ 05	44.50							
S10,	.23							
T10,	.04							
A1203	.32							
FeO MnO MgO CaO Na ₂ O K ₂ O Y ₂ O ₃ ^b C1 ^b	.52 .04 3.47 42.13 .31 .03 2.52 .02							
Total	94.13							
Additional detectio	ns ^C							
Cerium oxide (Ce ₂ 0 ₂)	2.0							
Neodymium oxide (Nd ₂ O ₂)	1.5							
Gadolinium oxide (Gd ₂ O ₂)	.5							
D;sprosium oxide (Dy ₂ O ₁)	.5							
Erbium oxide (Er ₂ 0 ₃)	.5							
Lanthanum oxide (La203)	.8							

TABLE 4-IV. - WHITLOCKITE IN TROCTOLITE C. AST

^aMean of six analyses. ^bY₂O₃ = yttrium oxide, Cl = chlorine.

^CApproximate values.

Gooley et al. (ref. 4-9) describe a troctolite of similar bulk composition, having slightly higher magnesian olivine content (Fo₈₈), in which the discrete spinel grains are compositionally variable with lower TiO₂ and aluminum oxide (Al₂O₃), higher chromium oxide (Cr₂O₃), and a lower magnesium value (Mg₄₅); titanum-rich phases such as ilmenite and armalcolite are absent, and whitlockite is an extremely rare accessory mineral. The symplectic intergrowth of diopside, enstatite, and spinel observed by Gooley et al. is not present in troctolite fragment 8A, and the spinel is an equilibrium phase associated with olivine and anorthite.

Roedder and Weiblen (ref. 4-10) describe a spinel troctolite fragment from Luna 20 soil samples having similar bulk composition, but the olivine in this sample is Fo_{91} and the spinel (Mg_{82}) contains 70 percent Al_2O_3 and 1.7 percent Cr_2O_3 . This sample has an igneous, quenched texture. In an earlier report, Roedder and Weiblen (ref. 4-11) described a slightly shocked olivine anorthosite (sample 14.30.3.51) containing spinel (Mg_{53}) of very similar composition to those in troctolite clast 8A with 18 percent Al_2O_3 and 46 percent Cr_2O_3 , but these spinels coexist with olivine of composition Fo_{74} to Fo_{77} .

LUNAR SAMPLE STUDIES

Compound or	Basalt	clast 4A	Tr	coctolite cla	Glassy basaltic					
e lemont	Spinel	Ilmenite	Spinel	Ilmenite	Armalcolite	fragment spinel				
Compound abundance, wt.%										
Si0 ₂	0.14	<0.03	0.12	0.07	0.03	<0.09				
TIO2	4.28	52.98	1.96	57.16	74.13	5.50				
Zr02 ^a	 .		<.04	<.04	.52					
A12 ⁰ 3	16.22	•0 9	25.54	.08 _	1.74-	15.60 _				
v203 ^a	.96	<.03	.27	.03	<.05					
Cr ₂ 0 ₃	40.24	.40	41_92	.30	1.20	42.14				
Fe0	32.76	43.55	18-31	31.39	11.51	33.02				
MnO	. 30	.39	. 19	. 32	.03	.26				
MgO	4.22	2.47	11.69	10_62	10.34	3.19				
Ca0	.15	. 12	.13	.10	.11	.22				
Total	99.27	100.00	100.13	100.07	99 . 61	99.93				
No. grains	1	4	6	7	3	1				
		Mol	lecular pro	oportion						
0	4.000	6.000	4.000	6.000	5.000	4.000				
Si	.005	0	.004	.003	.001	0				
Ti	. 109	1.978	.045	1.998	1.965	.139				
Zr			۹_	.0	.009					
Al	.646	.005	.916	.004	.072	.620				
V	.026	0	.007	.001	0					
Cr	1.075	.016	1.009	.011	.033	1.123				
Fe	.926	1.808	. 466	1.220	.339	. 931				
Mn	.009	.016	.005	.013	.001	.007				
Mg	.213	.183	.530	.734	.543	.160				
Ca	.005	.006_	.004 -	.005	.004	. 008				
Total cations	3.014	4.012	2.986	3.989	2.967	2.988				
Ratio ^b	18.7	9.2	53.1	37.6	61.5	14.7				

TABLE 4-V	OPAQUE	ANALYSES
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^a $2ro_2$ = zirconium dioxide, V_2o_3 = vanadium oxide.

^ບ100 Mg/Mg+Fe.

The large partition coefficient for Mg and iron (Fe) for coexisting olivine and spinel in sample 8A contrasts with much smaller partition coefficients in the two other "igneous" samples quoted. From this difference and from the very strongly selective partition against TiO_2 and zirconium dioxide (ZrO_2) in spinel and against

 ZrO_2 and Cr_2O_3 in illucente, the authors infer that the troctolite mineral assemblage equilibrated at comparatively low temperatures, well below the solidus but possibly above the temperature of equilibration of olivine and spinel and that of reaction between olivine and plagioclase in the troctolite described by Gooley et al. (ref. 4-9).

TROCTOLITIC AND BASALTIC CLASTS FROM A FRA MAURO BRECCIA



Figure 4-1.- Reflected light photomicrograph showing the association of spinel (Sp), ilmenite (Ilm), and armalcolite (Arm) in clast 8A; the stringers and small inclusions within the ilmenite are spinel.

The texture of the troctolite clast is shown in figure 4-2. Brown et al. (ref. 4-12) have observed 2- to 3-mm-size fragments of olivine (Fo₈₆) and plagioclase (An₉₅) intergrowths in breccia sample 14320,4 and describe their texture as crešcumulate, similar to the layered ultra-basic rocks of Rhum (ref. 4-13). The long, branching olivines characteristic of crescumulates are absent in clast 8A; olivine occurs as chains of randomly oriented, interlocking crystals set in a matrix of polycrystalline plagioclase. These chains could be remnants of a coarser, crescumulate texture, but, if so, recrystallization has been completed. Alternatively, deformations followed by complete annealing could produce the observed texture.

A glassy fragment fused to the troctolite clast along a 2-mm boundary in the thin section seems to be partly remeited mare basalt. This fragment contains. plagioclase, spinel, and pigeonite (tables 4-III, 4-V, and 4-VI) typical of such



Figure 4-2.- Ordinary light photomicrograph of troctolite clast 8A showing polycrystalline aggregates of olivine and plagloclase.

TABLE 4-VI.- PYROXENE ANALYSES

	-		<u> </u>	1				-		
Compound,	Ba	salt clast	4A	B	asalt clast (6A	Contact zone,	Glassy basalt,		
or mineral	Pigeonite	Mg-augite	Fe-augite	Pigeonite	Mg-augite	Fe-augite		precourte		
Compound abundance, wt.%										
Si0 ₂	53.01	50.53	48,60	52.67	50.45	50.11	56.61	52.39		
T102	.46	1.12	1.37	.55	1.09	1.62	. 35	. 61		
A1203	1.79	1.50	.26	1.92	3.45	1.91	1.59	1.46		
FeO MnO MgO CaO	15.99 .22 23.35 4.24	22.54 1 ³³ 14 ₁ 06 9.23	29.89 143 4.64 12.53	16.76 .31 22.57 4.91	15.76 .31 19.00 8.86	22.56 .40 12.98 10.64	9.99 .11 30.54 .49	18.22 .26 21.15 5.07		
Na20	<.04	<.04	.07	<.04	< 04	.06	<.04	<.04		
Cr203	.84	. 3 8	.06	.86	1.26	.56	. 31	.64		
Total	99÷90	99 . 69	98.8 5	100.55	100.18	100.84	99.99	99.80		
			Mol	lecular propo	ortion ^a			•		
Si	1.942	1.946	1.979	1.930	1.876	1.919	1,982	1.947		
Al ^{IV}	.058	.054	.021	.070	.124	.081	.018	. 053		
AI ^{VI}	.019	.014	,040	.013	,027	.005	.048	.011		
Ti	.013	. 032	042	.015	031	.047	.009	.017		
Fe	.490	.72 6	1.018	.514	490	.722	.292	. 566		
.Yn	.007	.011	1015	.010	.010	.013	.003	. 008		
Mg	1.275	.307	J282	1.234	1.053	.741	1.593	1.172		
Ca	. 166	. 38 1	.547	. 193	. 353	.437	.018	. 202		
Na	0	0	.0 06	0	0	.005	0	0		
Cr	.024	.012	.002	.025	.037	.017	,009	. 019		
Total	3.994	3.983	3.952	4.004	4.001	3.987	3.972	3.995		
			Miner	al abundance	e, mol%					
Enstatite Ferrosilite Wollastonite	66.0 25.4 8.6	42.2 37.9 19.9	15.3 55.1 29.6	63.6 26.5 9.9	55.6 25.8 18.6	29.0 38.0 23.0	83.7 15.4 .9	60.4 29.2 10.4		

^aOn the basis of six oxygen ions.

56

LUNAR SAMPLE STUDIES

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TROCTOLITIC AND BASALTIC CLASTS FROM A FRA MAURO BRECCIA

basalts, although an olivine grain, unusual in containing neither calcium (Ca) nor chromium, is also included. (See table 4-II for limits of detection.) Troctolite minerals at the boundary have interacted with the glassy basalt. Contacting olivine grains are strongly zoned. One large grain of typical Fo₈₆ composition adjacent to the troctolite plagioclase is zoned to Fo₅₆ at the glassy contact. A few homogeneous enstatité (En) grains (table 4-VI) are formed only along this contact region. Plagioclase, however, is not altered. These reactions are confined to the contact zone and do not influence the center of the clast. The reactions could have been caused by an event affecting the fragment edge before basait attachment or, more probably, by "contact metamorphism" produced by the basalt glass coating. Mineral grains in the glassy basalt are not zoneu, but available data are not sufficient to determine any possible annealing history.

The troctolite clast preserves evidence of an early low-temperature recrystallization and equilibration and a later contact with a fused but rapidly chilled basalt, which did not cause penetrative reequilibration of the troctolite. The troctolite clast was finally incorporated into the light matrix of the breccia. Grieve et al. (ref. 4-3) and Duncan et al. (ref. 4-14) recognize three generations of recrystallization in fragments within the breccia, the first at a mean temperature of approximately 1273 K (1000° C) and the second and third at 973 K. (700° C). This thermal history does not fit neatly with that of the troctolite, although the lower temperature could correspond to that of equilibration.

BASALT CLASTS 4A AND 6A

Basalt clasts 4A and 6A of sample 14321,88 are rounded fragments containing grains larger than 200 μ m. Clast 4A contains approximately 3 percent olivine, 5 percent opaque minerals, and more pyroxene than plagioclase, whereas clast 6A contains approximately 20 percent olivine and equal quantities of plagioclase and pyroxene with fine-grained ilmenite. Only one olivine grain was successfully extracted from clast 4A (table 4-II), and the only spinel (table 4-V) analyzed was included in this grain. Because the entire small, high-density separate of clast 6A was required for Rb/Sr dating, no ilnicinite analyses were obtained for this clast. The pyroxene data for both fragments are summarized in figure 4-3, and représentative analyses are given in table 4-VI. Average plagioclase analyses are given in table 4-III.

The olivine in clast 4A is considerably richer in Fe than the early pigeonite (fig. 4-3); consequently, the included spinel, although obviously a primary phase may not represent the initial chromite composition that crystallized. Nevertheless, if the titanium partitioning is comparable tothat in Apollo 12 basalts, then the titanium content of the spinel indicates that the TiO_2 content of the magma was approximately 3 percent. The pyroxene zoning of clast 4A is similar to that of the type VI lunar basalt classification of Bence and Papike (ref. 4.15). The representative of the type is sample 14053, an aluminous rock having a very low oxygen fugacity in the late stages of crystallization and apparently itself once a clast in a breccia. The spinel analysis may be matched precisely by primary nonreduced 14053 spinels analyzed in the authors' laboratory; however, the 14053 olivines are much more magnesian (Fo_{67} to Fo_{69}) and the feldspars more calcic (An₈₄ to An₉₄). Pyroxenes composed of 35 mol% wollastonite (Wo35) and En45 found in sample 14053 are absent in the specimen removed from clast 4A. Nevertheless, the evidence is that the clast has a bulk composition and an



Figure 4-3.- Pyroxene zoning in basaltic clasts 4A (crosses) and 6A (dots). Core and rim (greater ferrosilite content) analyses are joined.

LUNAR SAMPLE STUDIES

initial crystallization history sintilar to those of sample 14053 but continued to precipitate olivine to lower températures. Late-stage pyroxenes are zoned toward hedenbergite and were precipitated together with ilmenite. Unfortunately. no data exist for the final stages, which are characterized in sample 14053 by very low oxygen fugacities and recorded_by subsolidus_oxide reduction reractions.

The olivine in clast 6A is of similar Mg/Fe content to the most magnesian pigconite, and the early subcalcic augite contains as much as 3.5 percent Al_2O_3 . Apart from the Al_2O_3 content, the chemical trends in the pyroxenes are similar to those in clast. 4A and sample 14053, although zoning in the pyroxene quadrilateral does not extend as far in the hedenbergite direction. Clast 6A seems to be another high- Al_2O_3 basalt preserving minerals of a crystallization temperature earlier than those of thinerals in clast 4A.

RELATIONSHIP OF TROCTOLITE CLAST TO OTHER LUNAR HIGHLAND COMPOSITIONS

Prinz et al. (ref. 4-8) have described, analyzed, and classified 157 lithic fragments from the luttar highland region sampled by the Luna 20 spacecraft. The rock type classified as spinel troctolite has an average composition close to that of the troctolite 8A clast (table 4-I). The mineralogy given for spinel troctolite (table 2 and figs. 8 to 13 of ref. 4-8) shows olivine averaging Fo₈₃ and plagio lase averaging An₉₆ but spinel typically with more than 60 percent Al₂O₃ and less than 5 percent Cr2O3. Prinz et al. noted that, in bulk composition and mineralogy, the spiriel troctolite fragments form a tightly clustered group. Troctolite clast 8A therefore assumes a greater significance in exemplifying the subsolidus equilibrated mineralogy of this bulk compositional grouping. The discussions in references 4-8 and 4-16 of crystallization in the Fo-rich olivine/anorthite/silica system illustrate that the spinel troctolite bulk composition will crystallize as an equilibrated assemblage of olivine and anorthite plus accessory minerals, including chromium-rich spinel and minor orthopyroxene. However, if this assemblage is partly or completely melted in an impact event, the mineralogy will contain highly aluminous spinel as a major phase. Thus, the principal differences between the mineralogy of troctolite 8A and that of the spinel-bearing troctolites described in references 4-8 and 4-9 are due to differences in temperature of crystallization.

The primary origin of troctolite clast 8A may have been as a cumulate from magnesian, high- Al_2O_3 basalts of the "Fra Mauro type" (ref. 4-17). If this origin is valid, the presence of accessory titanium-rich minerals, whitlockite, and barian_K-feldspar establishes a link with the geochemical characteristics of the alkali-rich members of this basalt suite (e.g., high-K Fra Mauro basalts; potassium, rare-earth elements, and phosphorus (KREEP)). The cumulus origin would imply interstitial entrapment of source parent liquid and equilibration of this unit with the cumulus phases at_subsolidus temperatures.

An alternative to the preceding genetic relationship is to interpret the Fra Mauro basalts as having been formed by varying degrees of incomplete melting of a parent froctolite composition. Results of experimental studies (refs. 4-16 and 4-18) show that olivine (Fo₈₀ for KREEP, Fo₈₈ for sample 14310) and plagioclase are the nearliquidus phases for "Fra Mauro basalt" types at low pressure and would be the residual phases during partial melting. Incomplete melting of the troctolite would leave olivine (Fo_{86.5+}) and plagioclase (An_{95.5+}) as residual phases and eliminate phases such as barian K-feldspar, whitlockite, and ilmenite. Comparison of REE data for troctolite clast 8A and Fra Mauro basalts (ref. 4-4) shows that REE data, applied with current uncertainties in partition coefficients and assuming partial melting or crystallization in equilibrium, are consistent with either interpretation.

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U-Th-Pb Systematics of Apollo 16 Samples 60018, 60025, and 64435; and the Continuing Problem of Terrestrial Pb Contamination of Lunar Samples

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Uranium, thorium, and lead concentrations and lead isotopic compositions of Apollo 16 breccias 60018 and 64435 and anorthosite 60025 were determined. Serious terrestrial lead contamination of most 60018 separates and of sample 60025 occurred before sample digestion, although no uranium or thorium contamination was detected. The data from essentially uncontaminated breccia 64435, breccia 60018, and a glass separate of breccia 60018 yield uraniumthorium-lead pattern's much like those that the authors have found and published for other Apollo 16 rocks. An acid-leach experiment of one of the highly contaminated samples (< 100 mesh, 60018) revealed that the contaminate lead was not easily leachable and that it may have occurred by addition of noteasily-dissolved solids. Analyses of a "clean" NASA saw blade revealed 10 to 11 ppm lead in the saw's cutting edge, which may have been the source of lead contamination.

The lead (Pb) isotopic compositions of Pb and the concentrations of uranium (U), thorium (Th), and Pb were determined in Apollo 16 anorthosite 60025 and breccias 60018 and 64435, collected in the Descartes mountain region in the central lunar highlands.

Unfortunately, terrestrial Pb contamination grossly altered the Pb concentrations and U/Pb ratios of many of these samples. An effort was made to discover the source of this contamination.

The relatively uncontaminated samples supplement U-Th-Pb Apollo 16 data previously reported and augment U-Th-Pb patterns already established (refs. 5-1 and 5-2).

ANALYTICAL PROCEDURE

All whole-rock samples were crushed in a stainless steel mortat and analyzed directly. Phase separates of breccia 60018 were obtained by handpicking, sieving, and using a hand magnet.

Samples were dissolved in Teflon bombs. Extraction of the Pb was achieved using a two-step anion exchange technique (ref. 5-1). Total Pb blanks ranged from about 0.5 to 1.9 ng during this study. The U and Th were separated with Dowex 1×8 . NO₃-form resin (ref. 5-3). The U and Th blanks were less than 0.01 ng.

RESULTS AND DISCUSSION

Concentrations of U, Th, and Pb and the Pb isotopic compositions of breceias 60018 and 64435 and anorthosite 60025 are listed in tables 5-I and 5-II, respectively. The U concentrations in breceia 60018 range from ~ 0.1 ppm (plagioclase concentrate) to ~ 1.2 ppm (light gray lithic fraction) and probably

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LUNAR SAMPLE STUDIES

reflect varying amounts of potassium, rarc-earth elements, and phosphorus (KRFEP)¹ component. Plagioclase-pyroxene breccia 64435 contains only about 0.02 ppm U and 0.09 ppm Th – lower than all lunar rocks yet analyzed excluding anorthosites.

Data in tables 5-I and 5-11 are graphically displayed on a 206Pb/207Pb versus 238U/207Pb positive slope variant of the 207Pb/206Pb versus 238U/206Pb plot (fig. 5-1) of reference 5-4. Also plotted for comparison are breccia 66095 (ref. 5-2), hornfels 65015

¹Term used to express chemical composition of lunar materials. After Hubbard, N. J.; and Gast, P. W.: Chemical Composition and Origin of Nonmare Lunar Basalts. Proceedings of the Second Lunar Science Conference, vol. 2, MIT Press (Cambridge, Mass.), 1971, p. 999. (refs. 5-1 and 5-9), anorthosites 60015 (ref. 5-1), 15415 (ref. 5-6), and 67075 (ref. 5-8.), and another analysis of 60025 (ref. 5-5). Except for anorthosites 60015, 67075, and 60025, all Apollo 16 whole-rock samples previously published (refs. 5-1 5-2, 5-8, and 5-10), fall within error on the solid line drawn in figure 5-1. Chipped samples of hornfels 65015 with a 206Pb/204Pb ratio of >2500 (refs. 5-1 and 5-9) and breccia 66095 with a Pb content of \sim 15 ppm (ref. 5-2) represent pristine, essentially uncontaminated lunar material. The dashed contamination line in figure 5-1 defines the line along which hornfels 65015 would move if mixed with modern terrestrial Pb. The proximity of anorthosite 60025 and five of the separates of breccia 60018 to this contamination line and the rather low ²⁰⁶Pb/²⁰⁴Pb values of these samples (table 5-11) suggest that up to 99 percent of the

Sample	Run	Weight,	Cor	ncentrations,	Atomic ratios			
		8	υ	 Th	РЪ	232 _{Th} 238 _U	238 _U 204 _{РЬ}	
Anorthosite 60025	1 ^a 2	74.4 113.7	0.000594 .000502	0.00152 .00108	0.0652	2.64 2.22	0.581	
Breccia 60018:								
Whole rock	^a 1	208.9	.801	3.160	1_722	4.08	1378	
<100 mesh	esh 1 a ₂		.796 .381	3.021 1.398	6.643 1.369	3.92 3.79	9.55 190	
Handpicked glass	a ₁	32.3	.694	2.588	1.827	3.86	353	
Magnetics	1	29.8	.433	1.544	1.657	3.68	34.9	
Handpicked dark gray	1	39.6	1.055	3.584	2.623	3.51	161	
Handpicked light gray	^a 1	88.4	1.209	4.469	2.520	3.82	230	
Handpicked white	1	19.11	.1123	.544	.710	5.01	19.4	
Breccia 64435	1	58.4	.0226	.0917	.354	4.19	39.2	
	^a 2	145.6	.0299		. 397		35.6	

TABLE 5-1.- CONCENTRATIONS U, Th, AND Pb IN APOLLO 16 ANORTHOSITE 60025 AND BRECCIAS 60018 AND 64435

^aSamples totally spiked before digestion.

U-Th-Pb SYSTEMATICS OF APOLLO 16 SAMPLES 60018, 60025, AND 64435

Pb extracted from these samples is terrestrial. Breccia 64435, breccia 60018, and handpicked glass samples of breccia 60018, on the other hand, lie very close to or on the solid line int figure 5-1; have high 2^{06} Pb/ 204 Pb ratios (table 5-II); and appear to be negligibly contaminated with terrestrial Pb.

Minimally contaminated samples

When corrected for an assumed primordial Pb (ref. 5-11), the whole-rock and glass analyses of breccia 60018 fall within error on the 4.47- to 3.99-billionyears (b.y.) discordia line defined by other Apollo 16

Sample	Weight, mg	Run	Atomic ratios										
			0	bserved rat	los ^a	Corrected for analytical blank ^{b,c}							
			$\frac{\frac{206}{Pb}}{204}$	207 _{РЬ} 204 _{РЬ}	208 _{РЬ} 204 _{РЬ}	206 _{РЬ} 204 _{РЬ}	207 _{РЬ} 204 _{РЬ}	208 _{РЬ} 204 _{РЬ}	207 _{РЬ} 206 _{РЪ}	208 16 206 _{рь}			
••••••		•		Anort	iosite 6002	.							
Whole rock	95.6 74.4	P1 C1 d	18.52 18.73	15.72 15.71	38.14	18,67 18,83	15.78 15.74	38.37	0.8456	2.056			
	113.7		18.52	15.64	l	18.54	15.65	1	.8440	1			
······································		т · -		Bree	cda 60018	-	1	1		T			
Whole rock	56.9 208.9	р1 ^d с1	436.9 1119	244.3 606.0	432.0	643.2 1356	357.1 732.9	627.7 	0,5552	0,976			
(100 mesh (magnetics removed)	49.3 48.7 180.5	Р1 С1 d _{C2}	25.92 25.94 224.2	19.58 19.59 124.2	45.35 	26.02 26.00 239.0	19.65 19.62 132.0	45.58	. 7551 . 7546 . 5523	1.752			
Handpicked glass	71.9 32.3	P1 d _{C1}	262.3	144.1 219.6	272.2	298.2	163.0 274.2	307.4	.5466 .6762	1.031			
Magnet1cs	32.5 29.8	P1 C1	51.37 50.15	32.54 31.71	71.41	52.12 50.82	32.94 32.04	72.35	.6319 .6306	1,388			
Handpicked dark grav	40.9 39.6	14 1	148.1 169.7	77.68 88.16	169.0	151.7 174.2	74.60 90,32	173.0	.5248 .5185	1,141			
Handpicked light grav	109.2 88.4	P1 d _{C1}	230.2 216.8	110,2 104,3	<u>1</u> 33.2	247.2 227.1	117.8 106.8	244.5	. 4765 . 4769	1, (39			
Handpicked white	40,37 19,11	P1 	42.88 41.89	29.32 28.67	66,72	43.78 43.53	29,84 29,58	67.46	,6816 ,6795	1.552			
				Bre	octa 644 15								
Whole reck	71.0 58.4	P1 C1	159.9 150.5	148.6 186.1	152.8	223.5 221.1	280.8 277.4	204.8	1				
	145.6	°c2	155.2	195.3		170.6	215.4		1.263	· · ·			

TABLE 5-11.- ISOTOPIC COMPOSITION OF Pb IN APOLLO 16 SAMPLES 69025, 60018, AND 64435

^a208_{Pb} spike contribution subtracted from Pb concentration data.

The Pb (solupic composition used for blank corrections is $\frac{206}{Pb}/\frac{204}{Pb} \approx 18.06$; $\frac{207}{Pb}/\frac{204}{Pb} \approx 19.08$;

and 208pb/204pb = 37.88.

 $^{\circ}$ Analytical total Pb blanks were 1.18 ne for anorthostre 60025 analyses and ranged from 0.48 ne to 1.9 ne to 5 \pm remaining analyses.

dSamples totally spiked before direction.

63

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Figure 5-1.- A plot of 206Pb/207Pb compared to 238U/206Pb. New data from 60018, 60025, and 64435 are plotted. The following data are plotted for comparison: 60025 analyzed in reference 5-5, an interior analysis of anorthosite 60015 (ref. 5-1), breccia 66095 (ref. 5-2), anorthosite 15415 (refs. 5-6 and 5-7), anorthosite 67075 (ref. 5-8), and hornfels 65015 (ref. 5-1). Data are corrected for blanks only.

whole-rock U-Pb data excluding anorthosites (refs. 5-1 and 5-9). The 60018 whole-rock analysis is nearly concordant at about 4.42 b.y.² The glass contains excess Pb relative to U, which suggests it is either a soil impact melt (i.e., the excess Pb may have been present before glass formation) or the glass somehow gained Pb relative to U after its formation.

Although breecia 64435 contains only ~ 0.4 ppm Pb, its very low U and Th contents indicate a severe

net enrichment of Pb relative to U and Th. Much like breccia 66095 (ref. 5-2), breccia 64435 apparently gained Pb characterized by a very high 207Pb/206Pb ratio of about 1.45. The isotopic character of the excess Pb in 64435 requires that this Pb was produced in a U-rich reservoir very early in the Moon's history when the production rate of 207Pb exceeded that of 206Pb. Model ages of 3.73 to 4.0 b.y. for the introduction of excess Pb into breccia 64435 and of 4.65 to 4.42 b.y. for the production of a U-rich reservoir ($\mu = 496$ to 1115) may be calculated using twoand three-stage U-Th-Pb evolution models, respectively, as discussed at length in reference 5-2.

Despite gross differences in absolute U. Th. and Pb concentrations measured in breccias 64435 (table 5-1)

²The constants used are $\lambda_{238} = 0.15369 \times 10^{-9} \text{ yr}^1$; $\lambda_{235} = 0.97216 \times 10^{-9} \text{ yr}^{-1}$ (ref. 5-12); and 238U/235U = 137.8 (ref. 5-13). If the U-decay constants of reference 5-14 were used, the ages reported here would be reduced approximately 1.3 percent.

U-Th-Pb SYSTEMATICS OF APOLLO 16 SAMPLES 60018, 60025, AND 64435

and 66095 (tef. 5-2), both these samples appear to have undergone very similar U-Th-Pb evolution histories.

The Continuing Problem of Terrestrial Pb Contamination of Lunar Samples

Serious terrestrial Pb contamination of some of the Apollo 16 samples has occurred. In this section, the cause of contamination is discussed, and procedures whereby such contamination may be eliminated in the future are recommended.

Contamination of lunar samples by terrestrial Pb has long been a problem to U-Th-Pb investigators. Possible sources of contamination discussed in the literature are as follows.

1. Pb introduced by analytical blanks (currently only a serious problem in very small samples or in samples with very low Pb concentrations such as anorthosites)

2. Mineral-separation techniques (e.g., the heavyliquid problem reported in ref. 5-7)

3. NASA preparation of the samples (e.g., the contaminated sawdust reported in refs. 5-7 and 5-15)

4. Rocket exhaust contamination possibilities mentioned in reference 5-7

5. Contamination on the lunar surface during sample collection (e.g., the core tube contamination of Apollo 15 and 16 samples reported in refs. 5-8 and 5-16)

Handpicked mineral concentrates and sieve fractions of breccia 60018 and whole-rock analyses of 60025 show strong apparent terrestrial Pb contamination (table 5-III). In particular, the Pb isotopic composition of 60025 determined in the authors' laboratory was indistinguishable from that of modern terrestrial Pb (206 Pb/ 204 Pb \sim 18.5; measured Pb concentrations = 0.065 and 0.094 ppm). One split of the <100 mesh fraction of 60018 contained 6.6 ppm Pb with a 206 Pb/ 204 Pb ratio of 26.01; whereas relatively uncontaminated whole-rock and glass samples from this same rock suggest this ratio should be at least 10 times this value³. Altogether, three separate splits of the <100 mesh fraction (minus the magnetic component) were analyzed. Two of these splits were analyzed directly (tables 5-I and 5-II), and one was used for an acid-leach experiment (table 5-IV). The gross differences in the measured U, Th, and Pb concentrations and Pb isotope ratios of these three samples reflect considerable heterogeneity among the splits. The acid-leach experiment sliowed that the residue contained as much Pb as the preceding three acid leaches combined and had a lower $^{206}Pb/^{204}Pb$ ratio than any of the acid leaches (table 5-IV). This finding probably indicates a particulate type of contamination rather than easily dissolved surface contamination.

. Interestingly, the Th/U ratios found in the first two nitric acid leaches (steps 2 and 3, table 5-IV) are much higher than the third acid leach (step 4, table 5-IV), which in turn has a lower Th/U ratio than does the residue (step 5, table 5-IV). This pattern was also found in the leach experiment of breccia 66095 (ref. 5-2), and shows that Th is bound in at least two very different sites. Possibly the nitric acid leaches dissolved one or more accessory Th-rich phases such as phosphates.

Because sample 60015 (ref. 5-1) falls outside of error off the U-Pb trend established for other Apollo 16 rock samples (fig. 5-1) in the direction of modern terrestrial Pb, this sample may also exhibit a small degree of terrestrial Pb contamination. Alternatively, this sample may contain a lunar Pb distinctly different from that of most other Apollo 16 samples (ref. 5-1).

The Pb data for 60025 (this report and ref. 5-5) are not sufficiently precise, owing to terrestrial contamination problems, to calculate model ages – rather, these data emphasize that some lunar plagioclase may contain almost nc Pb at all.

The large quantity of Pb measured in several of the contaminated samples (especially the first <100 mesh split of 60018 with 6.6 ppm Pb) is impossible to explain by physical or analytical contamination in the authors' laboratory. All samples analyzed by the authors that exhibit significant contamination have one thing in common: they were cut with a bandsaw in the NASA Lunar Receiving Laboratory, although not all samples so dissected were significantly contaminated (table 5-III). To pursue this problem further, the authors requested and received a "clean" saw blade from NASA that had been cleaned as re-

³The Pb concentration determinations were made from 10.6 ng and 4.9 ng of total Pb extracted in the case of 60025 and 324 ng Pb extracted in the case of 60018 <100 mesh split 1. Analytical blanks associated with these 6:025 and 60018 analyses were 1.18 ng and 1.72 ng, respectively.

TABLE 5-III. - APOLLO 16 SAMPLES ANALYZED BY NUNES AND TATSUMOTO

[Percentage lead contamination estimated by Nunes and Tatsumoto (ref. 5-1) from the degree analyses depart from the U-Pb trend defined by a line joining 66095 and 65015 whole-rock data]

Sample no. Weight, A g c		Apparent l⊬ad contamination, percent	Any sawed surface? ^a	Comments ^a				
60015,46	6.83	13	Yes	Thin chips with sawed surfaces along the narrow edges.				
60018,16: 100 mesh Handpicked plagioclase Magnetic fraction Handpicked dark gray Handpicked light gray Handpicked glass Handpicked whole rock	21.67	77 64 53 32 12 (b) (b)	Yes, on almost all faces	Piece of 1.5-cm-thick slab; sufficient material existed in the slab so that sawed surface could be chiseled off to produce 5-g pieces free of sawed surface.				
60025,9003	2.81 99		Yes, on about 60% of faces	Sample is all of ,26 furnished by Haskin.				
60315,81	3.04	(ь)	No	Fresh chips (no sawed or lunar exterior surfaces).				
64435,5-	10	(b)	Some	Sample obtained from Rosholt; four large interior chips.				
65015,52	3.09	(c)	No	Fresh chip.				
66095,41	4.85	(c)	Ne	Rock was not sawed.				
67015,11	5.08	(Ь)	No	Rock was not sawed.				
67015,12	4.96	(Ь)	No	Rock was not sawed.				
68415,60	3.48	(b)	~50% sawed surface					

^aSawed surface estimates and comments by P. Butler, Jr.

^b<2% terrestrial Pb contamination.

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Assumed to contain no terrestrial Pb.

66

LUNAR SAMPLE STUDIES

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Step	Leach description	Rum	Ръ	Ľ	Th	Atomic ratios ^d									
(6)		(6)	ng	ng	removed, Ng	Observed		Corrected for blank							
						²⁰⁶ РЬ ²⁰⁴ РЬ	²⁰⁷ ръ ²⁰⁴ Ръ	206 _{РЬ} 204 _{РЬ}	²⁰⁶ РЬ 204 _{РЬ}	207 _{РЬ} 204 _{РЬ}	²⁰⁸ рь ²⁰⁴ рь	207 Pb 200 Pb	206 _{Pb} 206 _{Pb}	2 12 ₁₆ 2 38 ₀	2.181 204 Pb
1	4 mi water; 4 days	P C	1.05			(23.62) (23.39)	(18.72) (18.25)	(48.16)	(27,21) (30,69)	(20.85) (22.14)	(55.35)	(0.7677) (.7214)	(2.034)	 	
2	<pre>4 ml 1 normal (18) nitric acid (EMO₃); 2 days</pre>	P C	157.2 158.6	46.43 46.87	509.9 515.1	44.66 44.66	29.95 29.95	89.'' 	44.31 ÷4.24	30.11 30.05	89.83	.6794 .6794	2.027	 11,35	42.1
3	4 mai 1639 BBEO ₃ ; 3 daaya	P C	254.6 296.0	26.99 31.45	218.4 254.6	51.44 51.50	37. 8 9 37.93	88.04 	51.70 51.64	38.07 38.02	88.62	.7364 .7364	1.714	 6. 16	 16.4
•4	2 ml 50% hydrofluoric acid (HF) + 2 ml 16N HENO ₃ ; 3 daym	P C	400.9 271.8	136 7 91.3	.958 .650	66.99 67.16	39.53 39.63	75.23	67.43 67.61	39.75 39.85	75.77 	. 5896 . 5895	1.124 	 .607	 53.3
5	4.5 ml 502 HF + 4.5 ml 1618 HDD ₃ ; haate? overmight (residue fiesolution)	P C	897.9 520.4	219.8 127.4	711.0 412.0	41.04 41.12	26.66 26.72	54.16	41.20 41.25	26.75 26.78	54.43 	.6493 .6493	1.321 ••	 3. 34	26.0
	Total		2960	726.9	2623	••									
	Apparent pps of total sample		2.97	.728	2.63					••					

TABLE 5-IV.- LEAD ISOTOPIC COMPOSITIONS AND C, TH, AND PE CONTENTS OF ACID LEACHES AND RESIDUE OF A THE MESH SAMPLE" (MAUNETICS REMOVED) OF BREACIA 65018

The total sample weighed 995.7 mg; 208 Pb spike contributions were subtracted from concentration data.

b Total Pb blanks were 0.47 ng for leach steps 1, 2, and 3; 1.0 ng for step 4; and 1.7 ng for the step 5 residue dissolution.

^CP = composition. C = Concentration.

dParentheses indicate significant uncertainty because of the small amounts of lead present.

About one-third of the step 4 leach solution was spilled before dividing and spiking the concentration portion of the solution; thus, the totals of the 0. The and Pb contents are slightly lower than the true values.

4

LUNAR SAMPLE STUDIES



Figure 5-2.- Pieces of NASA steel bandsaw blade that were analyzed for Pb concentration (drawn approximately to scale).

quired by the NASA specification entitled "Lunar **Receiving Laboratory Cleaning Procedures for Con**tamination Control." Four pieces were cut from the blade: two pieces contained varying amounts of the weld but no cutting edge, one piece contained only nonwelded steel, and the fourth piece included some steel and the cutting edge (fig. 5-2). The saw blade santples were cut with an acid-cleaned pair of tinsnips, and all four samples were consecutively washed in twice-distilled acetone for about 10 minutes, in cold, 1.7 normal hydrochloric acid for about 5 minutes, and in five-times-distilled water to remove any surficial Pb. The saw blade samples were then dissolved in hydrochloric acid and nitric acid, and the total samples spiked. Pb separation was achieved with anion exchange columns.

The total Pb contents calculated assuming a 208Pb/206Pb sample ratio of 2.063 are shown in figure 5-2. Agreement of the Pb contents of the three saw blade samples with no cutting edge indicates the steel has a uniform Pb content of ~ 3 ppm regardless of the amount of welded region present. This value compares to the Pb content of 0.6 ppm found (ref. 5-15) for the copper-wire saw blade used in cutting Apollo 12 and 14 rocks.

CONCLUSIONS

The high Pb content of the bandsaw blade sample that includes a portion of the cutting edge (\sim 5 ppm) indicates Pb is concentrated in the cutting edge,

which consists of diamonds cemented in a white bonding material. It is emphasized that the measured Pb contents in these saw blade samples are minimal because of the preliminary acid-washing procedure. Judging from the relative amounts of steel and cutting edge in sample 4, a Pb content of ~ 10.8 ppm is estimated for the cutting edge material. This quantity of Pb is sufficient to explain all contaminated data. The variable amounts of Pb contamination (from insignificant to much) of bandsaw-cut samples may be explained as varying saw blade particulate contributions to the samples. This variance can explain, for example, why the <100 mesh fraction of 60018 was so highly contaminated (the starting material containing considerable saw-blade-cut surfaces); whereas the whole-rock sample displayed no obvious contamination, perphaps because it was picked from small chunks with no flat (cut) surfaces. Unfortunately, an accurate assessment of the amount of saw blade surface on any given sample dissolved was not made before the realization that these samples were contaminated. In any case, the particulate nature of the contamination, as indicated by the acid-leach experiment and the very high Pb content of at least 10 to 11 ppm found in the cutting edge of the bandsaw blade, indicates that significant Pb contamination of lunar samples during sawing was likely.

To ensure "clean" lunar samples in the future, the following procedures are suggested.

I. Lunar investigators working with Pb should request only chipped samples.

2. When only sawed material is available, chough material should be obtained so that individual investigators may themselves chip away material that displays no sawed surfaces.

3. Sample chunks of cohesive material that are derived from originally sawed samples should be washed in twice-distilled acetone and vibrated ultrasonically to remove any particulate material that may be foreign to the sample. This cleaning should be performed by the individual investigators before crushing and mineral separation.

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U-Th-Pb SYSTEMATICS OF APOLLO 16 SAMPLES 60018, 60025, AND 64435

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