A general review of lunar science is presented, utilizing two themes: a summary of fundamental problems relating to the composition, structure, and history of the moon and a discussion of some surprising, unanticipated results obtained from Apollo lunar science. (1) The moon has a crust of approximately 60-km thickness, probably composed of feldspar-rich rocks. Such rocks are exposed at the surface in the light-colored lunar highlands. Many highlands rocks are complex impact breccias, perhaps produced by large basin-forming impacts. Most highlands rocks have ages of \(\sim 3.9 \times 10^9\) yr; the record of igneous activity at older times is obscured by the intense bombardment. The impact rate decreased sharply at 3.8-3.9 \(\times 10^6\) yr ago. The impact basins were filled by flows of Fe- and, locally, Ti-rich volcanic rocks creating the dark mare regions and providing the strong visual color contrast of the moon, as viewed from earth. Crustal formation has produced enrichments in many elements, e.g., Ba, Sr, rare earths, and U, analogous to terrestrial crustal rocks. Compared with these elements, relatively volatile elements like Na, K, Rb, and Pb are highly depleted in the source regions for lunar surface rocks. These source regions were also separated from a metal phase, probably before being incorporated into the moon. The physical properties of the lunar mantle are compatible with mixtures of olivine and pyroxene, although Ca- and Al-rich compositions cannot be ruled out. Deeper regions, below \(\sim 1000\) km, are probably partially molten. (2) Lunar rocks cooled in the presence of a magnetic field very much stronger than the one that exists today, owing either to dynamo action in an ancient molten core or to an external magnetization of the moon. Lunar soil properties cannot be explained strictly by broken-up local rocks. Distant impacts throw in exotic material from other parts of the moon. About 1% of the soil appears to be of meteoritic origin. Vertical mixing by impacts is important; essentially all material sampled from lunar cores shows evidence of surface residence. The surface layers of lunar material exposed to space contain a chemical record of implanted solar material (rare gases, H) and constituents of a lunar atmosphere (\(^{40}\)Ar, Pb). Large isotopic fractionation effects for O, Si, S, and K are present. Physical properties of the surface layers are dominated by radiation damage effects. Lunar rocks have impact craters (\(\leq 1\) cm) produced by microgram-sized interplanetary particles. The contemporary micrometeorite flux may be much higher than is indicated by the microcrater densities, indicating time variations in the flux. Particle track studies on the returned Surveyor camera filter first showed that the Fe nuclei were preferentially enhanced in solar flares.

A. INTRODUCTION

Any flashback on the scientific accomplishments of the past 4-6 yr in the earth and planetary sciences must include a summary of Apollo lunar science. A priori it was probable that major advances in the understanding of the nature and history of a second planet would be forthcoming once lunar samples were available for laboratory study. This has in fact happened. Because of the vast amount of information available it is impossible for any one person to write a scientific summary of Apollo, even if he has lived through the excitement from the very beginning. Also, no two lunar scientists will probably agree on the relative importance of various discoveries. So with all due respect to my colleagues and without any pretense for completeness I want to reminisce on some of the findings of Apollo lunar science that I have found particularly intriguing. This paper is meant for other earth scientists who have not tried to follow the pace of lunar science.

If in the pre-Surveyor era I had written a realistic list of fundamental lunar problems that might be solved or at least significantly constrained by the opportunity to study lunar materials in terrestrial laboratories, the list of the important questions might have read as follows:

1. What are the differences in highlands and mare materials? Even to the naked eye the lunar surface is not homogeneous (Figure 1). From telescopic observation it was known that the dark (mare) regions were lower and less densely cratered than the lighter-colored highlands, which have a very high density of craters. The differences in crater density indicated that the mare surfaces were younger. Telescopic observations suggested that the mare were lava flows, but the basic distinction between the mare and the highlands rock types were unknown, except that the distinction was likely to be chemical in nature.

2. What is the composition of the moon as a planet? The nonuniform appearance of the moon suggested that planetary differentiation had occurred. But was this a predominant or a minor effect? Could inferences about the composition of the moon as a whole be drawn from chemical studies of returned samples?

3. Why is the density of the moon low, compared with that of the earth or the other inner planets? The large density contrast between the moon (3.34 g/cm³) and the earth (5.52 g/cm³) can only partially be accounted for by the self-compression effects of a comparatively large planet like the earth.
Fig. 1. Sketch of the moon as made by the Czech astronomer Andel in 1926. The Apollo and Luna landing sites are indicated. The dark, sparsely cratered areas (maria) are topographically low in comparison with the light-colored, densely cratered highlands. The Apollo 15 landing site is on the bottom right rim of the large impact basin, Mare Imbrium. Figure courtesy of G. P. Russ.

earth [Kovach and Anderson, 1965]. Moreover, the extreme comparison is between the moon and Mercury, which although its diameter is only 1.4 times the diameter of the moon, has a density of 5.5 g/cm³. It had been recognized for many years that these density differences must reflect chemical variations, even within the inner solar system, most reasonably reflecting the relative abundance of iron, the most abundant high atomic weight element.

4. What is the internal structure of the moon? The moment of inertia of the moon is consistent with that of a homogeneous planet [Tokšić et al., 1974]. But constrained only by the mass, total density, and moment of inertia, a large degree of internal structure is still permitted.

5. What is the state of evolution of the moon? Thermally, the earth is an active planet. The great majority of the surface and near-surface rocks of the earth have ages that are only a fraction of the age of the planet itself. In contrast, meteorites are primarily debris that has survived intact from the beginning of the solar system. Where does lunar material fit into this hierarchy?
Fundamental problems always turn out to be difficult; the above list is no exception. Furthermore, it is clear that the above problems cannot be isolated totally and attacked independently; they are strongly intertwined. It was not obvious, pre-Apollo, that any answers to these questions could be obtained, and completely satisfactory solutions on all counts do not exist at present. But is is important to reflect on where things stand and to consider what are the prospects for the future with respect to these major problems. This will be one theme of this paper.

However, there is a second theme that I wish to develop in parallel. The above problems could have been formulated without specific knowledge of lunar materials. In this sense they were 'anticipated' problems to be studied, even though the results obtained were in many ways surprising. However, in any venture into an unexplored research area, one always also hopes to find exciting 'unanticipated' results, and I believe that this was the case with lunar science. Even though it is possible that in some cases the results were unanticipated only by me, I found the following results new and exciting: (1) a highly nonuniform cratering rate in the early history of the moon, (2) impact breccias as the dominant highlands rock type, (3) a lunar paleomagnetic field, perhaps as large as that of the earth, (4) exotic components in the lunar soils (regolith), (5) fundamentally different properties for lunar surface layers, (6) microcraters on rocks, and (7) an enhanced solar flare iron flux.

For reasons of simplicity, brevity, and personal preference the discussion of the above two lists will focus on general interpretations and conclusions that are relatively 'model independent.' Many models have been formulated to address some of these topics more fully and to provide more specific conclusions than those given below. Such models are vital to lunar science. They are the quantitative manifestations of the qualitative observations that I will discuss. For the most part they are intelligently formulated, plausible, and perhaps in some cases even correct, but given the scope of this article, most discussion of these models will be by reference.

I make no pretense for complete referencing; all references cited are to be interpreted as having the implicit prefix, 'see, for example.'

B. FUNDAMENTAL LUNAR PROBLEMS

1. Mare Versus Highlands Rocks

The Surveyor missions showed that the lunar maria were composed of volcanic rocks [e.g., Turkevich, 1974], confirming earlier suggestions made by telescopic observations. Returned materials from Apollo 11 (Mare Tranquillitatis), Apollo 12 (Oceanus Procellarum), Apollo 15 (Mare Imbrium), Apollo 17 (Mare Serenitatis), and Luna 16 (Mare Fecunditatis) (Figure 1) have led to the establishment of a rock type that has become known as 'mare basalt' [Papike et al., 1974; Hays and Walker, 1975]. Figures 2–4 illustrate the distinctive chemical properties of mare basalts, as compared with compositions of highlands rocks and terrestrial basalt. Figure 1 shows that the much-publicized high Ti contents of mare basalts are confined to samples returned from Apollo 11 and 17. Telescopic observations indicate that these sites are among the darkest of the moon [Head, 1974]. If this correlation is generally valid, rocks of 10–15% TiO₂ are not rare but are probably somewhat atypical of the maria as a whole, although many relatively high Ti regions were seen by the orbital gamma experiment [Metzger et al., 1974]. The low Ti mare basalts are more similar to their terrestrial counterparts; however, essentially no terrestrial basalt is compositionally equivalent to the high Ti mare basalts. The existence of high Ti basalts is generally attributed to an enrichment in Ti for the moon as a whole and is paralleled by an analogous enrichment of many other such 'refractory' elements in lunar materials (see section B2). As Figures 3 and 4 show, mare basalts are also Fe-rich, alkali-poor rocks in comparison with terrestrial basalts. On the other hand, lunar and terrestrial surface rocks, as well as many achondritic meteorites, have much lower Fe/Si and Mg/Si ratios than chondrites; this can be viewed as a natural consequence of planetary magmatic differentiation processes in an Fe- and Mg-rich solar system.

The high Fe and Ti contents in mare basalts are expressed mineralogically as Fe-rich clinopyroxenes plus ilmenite and other Fe-Ti oxides, some previously unknown terrestrially [Haggerty, 1973]. The high abundance of these dark minerals and their relative absence in highland rocks account for the striking visual contrast of the near side of the moon. Although they are far from a homogeneous population, highlands rocks are all characterized by high proportions of Ca-rich plagioclase feldspar, reflected by high Al₂O₃ (Figure 1) and low FeO (Figure 3), and are consequently light-colored rocks. The Al-rich, Fe-poor nature of highlands rocks was first established by Surveyor 7 [Patterson et al., 1970]. Anomalous anorthositic fragments in the Apollo 11 soils [Wood et al., 1970], feldspar-rich rocks (including pure anorthosites) from

Fig. 2. Correlation diagram of Al₂O₃ versus TiO₂, showing fields occupied by mare basalts, highlands rocks, and chondrites. No lunar soils have been plotted. The insert compares the lunar fields with the one-standard-deviation spread in the distribution of Al₂O₃ and TiO₂ analyses for all terrestrial basalts, including both continental and oceanic samples, as compiled by Manson [1967]. The terrestrial range is approximate; any correlation of Al₂O₃ and TiO₂ has been neglected. The high Ti mare basalts are from Apollo 11 and 17 and differ greatly from any terrestrial counterpart. The low Ti mare basalts are from Apollo 12 and 15. Mare and highlands rocks are clearly distinguished on this plot. Lunar analyses are taken from papers by Rose and his co-workers, Rhodes and his co-workers, and Compton and his co-workers in the Lunar Science Conference proceedings.
I.0
0.8
0.4
0.2
0.0
0 0.5 1.0 1.5
K2O wgt %

Fig. 3. Correlation diagram of individual analyses of lunar rocks and an approximate field for terrestrial basalts defined by the standard deviation of the distribution of Fe and MgO analyses for basalts as compiled by Manson [1967]. Both oceanic and continental samples are represented. Lunar data are from sources given in caption of Figure 2. Mare and highlands rocks are clearly distinguished on the basis of their iron contents.

Apollo 14, Apollo 16, and Luna 20, plus the mountain front areas sampled during Apollo 15 and 17 provide our present sampling of highlands rocks (Figure 1). The overall Al-rich character of the lunar highlands has been established by uniformly high Al/Si ratios for all highlands areas observed by the orbital X ray fluorescence experiments on Apollo 15 and 16 [Adler et al., 1973].

As was indicated in the introduction, the high proportion of breccias returned from the highlands sites was a major surprise. Because of analogies to breccias from terrestrial impact craters and because of the domination of lunar surface features by the impact processes these rocks are universally believed to be impact breccias [James, 1975]. Some breccias are very weakly consolidated and contain distinct rock and mineral fragments. In some spectacular examples, e.g., 15465, the fragments are cemented together by thick veins of bubbly glass. Other breccias are much more coherent and uniform in appearance, presumably owing to a high degree of postimpact thermal annealing [James, 1975; Phinney et al., 1975]. Many petrologists believe that the bulk of the highlands breccias are the products of very large, possibly basin-forming impact events. A particularly strong case can be made for Apollo 14 breccias [Chao et al., 1972], because the landing site was on a ridge of ejecta extending southward (Figure 1) from the huge Imbrium impact basin. Many petrologists believe that some highlands rocks that show igneous textures (e.g., 68415 and 14310) are derived from impact-produced melts [Phinney et al., 1975].

2. Chemical Composition of the Moon

Figures 2–4 show that magmatic and impact processes have produced lunar materials of a wide variety of compositions. Looking through these effects would be impossible with only major element data; however, because of the large body of high-quality trace element and isotopic data on lunar samples, it is possible to make some qualitative inferences about the composition of the moon as a planet.

Lithophile element abundance patterns. Figure 5 shows the concentration ranges observed for six important elements expressed as a ratio of the lunar rock abundance to that for average solar system nonvolatile material. As is customary,

Fig. 4. Correlation diagram of Na2O versus K2O in lunar rocks (individual data points). The field described by the various chondrite classes is indicated. The insert is the same plot on a larger scale. The area enclosed by the dashed line indicates the average of various terrestrial basalt subtypes as defined by Manson [1967]. The terrestrial field may overrepresent alkali basalt types; nevertheless, the alkali-poor character (particularly for Na) of many lunar rocks is shown. The region south of Mare Imbrium is rich in K as shown by the Apollo 14 rocks. Lunar data are from sources given in caption of Figure 2.
lithophile, i.e., that show preferential incorporation into elements in lunar rocks relative to chondrites. Fields are indicated separately for low Ti mare basalt (Apollo 12 and 15 and 14053), high Ti mare basalt (Apollo 11 and 17), and 'highlands rocks' (Apollo 14 and 16 plus rocks sampled from mountain front stations on Apollo 15 and 17). Fields are indicated where there appears to be a continuum of data points, but if individual rocks can be plotted as recognizable discrete points, this has been done. Low abundance rocks, designated by triangles, are listed as anorthosites in the original literature sources. The discrete low Ti mare basalt samples with enhanced abundances are 12038 and 14053. The Apollo 11 'high K' basalts appear as a discrete blob at ~70 times chondrites for Ba and Th for the high Ti mare basalts. Data were obtained from papers by Philpotts and his co-workers, Gast and Hubbard and their co-workers, and Tatsumoto and his co-workers in the second volume of the various Lunar Science Conference proceedings, as well as from Tatsumoto [1970], and Tera et al. [1970]. The features of this plot are not sensitive to the class of chondrites used for reference. Chondritic abundances are from Philpotts et al. [1972] plus 9000 ppm for Na and 46 ppb for Th.

Fig. 5. Abundance patterns of lithophile elements in lunar rocks relative to chondrites. Fields are indicated separately for low Ti mare basalt (Apollo 12 and 15 and 14053), high Ti mare basalt (Apollo 11 and 17), and 'highlands rocks' (Apollo 14 and 16 plus rocks sampled from mountain front stations on Apollo 15 and 17). Fields are indicated where there appears to be a continuum of data points, but if individual rocks can be plotted as recognizable discrete points, this has been done. Low abundance rocks, designated by triangles, are listed as anorthosites in the original literature sources. The discrete low Ti mare basalt samples with enhanced abundances are 12038 and 14053. The Apollo 11 'high K' basalts appear as a discrete blob at ~70 times chondrites for Ba and Th for the high Ti mare basalts. Data were obtained from papers by Philpotts and his co-workers, Gast and Hubbard and their co-workers, and Tatsumoto and his co-workers in the second volume of the various Lunar Science Conference proceedings, as well as from Tatsumoto [1970], and Tera et al. [1970]. The features of this plot are not sensitive to the class of chondrites used for reference. Chondritic abundances are from Philpotts et al. [1972] plus 9000 ppm for Na and 46 ppb for Th.

There are at least two factors that contribute to the abundance pattern shown on Figure 5 and are also probably responsible for the tendency of lunar rocks to be Ti rich (Figure 2). These factors are (1) the formation of a lunar crust, presumably by internal differentiation of the moon, and (2) a systematic enhancement in the moon as a whole for those elements that are both refractory (highly nonvolatile) and lithophile, i.e., that show preferential incorporation into silicate phases (as opposed to sulfide or metal).

There is abundant chemical evidence for the existence of a lunar crust qualitatively analogous to the crust of the earth. The elements shown in Figure 5 also tend to be enriched in many terrestrial crustal rocks; however, this is only a suggestive argument. A much stronger argument follows from the Th and U concentrations in lunar samples. If the Th in the moon as a whole were as high (~0.5 ppm) as is measured in most lunar samples, then the moon as a whole would be molten today throughout most of its interior [Toksoz and Solomon, 1973]. This is shown not to be the case by seismic studies (see next section). Further, the lunar heat flow would be much higher than is actually observed [Langseth et al., 1973]. Thus the conclusion is inescapable that Th and other elements that have similar geochemical behavior have been differentiated upward into a lunar crust. Orbital gamma ray measurements have shown that rocks with ~0.5-ppm Th are not rare on the lunar surface but that rocks with very high (~3.5 ppm) Th tend to be concentrated on the lunar near side, particularly in the areas south of Mare Imbrium [Metzger et al., 1974]. Many such rocks were sampled by Apollo 14 and account for the highest enrichments (~100) shown on Figure 5 for all elements.

Thus to a first approximation, lunar chemistry is dominated by the crustal formation process, and this process must be dealt with before conclusions can be drawn about the nature of the source materials and the composition of the moon as a whole. The general types of data on which models of lunar magmatic processes can be based are: (1) experimental petrologic studies of the compositions of the liquidus phases for actual and hypothetical lunar bulk compositions [Hays and Walker, 1975] and (2) the relative abundances of trace elements (e.g., rare earths [Haskin et al., 1974; Weill et al., 1974]) or isotopes (e.g., initial $^{87}\text{Sr}/^{86}\text{Sr}$ [Papanastassiou and Wasserburg, 1972a]). The simplest cases to discuss are abundance ratios for elements that appear to be relatively invariant in magmatic processes. An important example is the K/U ratio, which as was pointed out by Wasserburg et al. [1964], is remarkably invariant at 1-2 x 10$^4$ for terrestrial igneous rocks and as is illustrated in Figure 6, has a value of ~2 x 10$^4$ for lunar rocks with K contents of 0.02-2%. The total range in K/U for both the lunar and the terrestrial samples shown is about plus or minus a factor of 2. Both the lunar and the terrestrial K/U ratios are distinctly lower than the ratio for ordinary chondrites, although the terrestrial and carbonaceous chondrite values are not clearly resolved. The correlated enhancement of K and U in the liquid phase until the final stages of crystallization of a mare basalt has been shown by U microdistribution studies [Burnett et al., 1971b].

The remarkable coherence of K/U implies that it can be considered a 'planetary constant' and that in particular, K/U ~ 2 x 10$^4$ is a general property of at least lunar crustal rocks and their source regions. Examples of other comparatively invariant lithophile element ratios are Zr/Hf [Ehmann and Chyi, 1974], Fe/Mn, K/Ba [Wänke et al., 1973], Th/U [Toksoz and Solomon, 1973], and Ti/U [Krähenbühl et al., 1973]. In the absence of the observed uniformity of the relative K/U/Th abundances, calculations of the thermal history of the moon and other planets [Toksoz and Johnston, 1975] would not be meaningful.

The low lunar K/U ratio compared with that for chondrites is generally interpreted as reflecting preferential enhancement of refractory elements in the moon and a corresponding depletion in elements that are even moderately volatile, e.g., K, Rb, and Cs. The volatility of Na is presumably also the reason for its apparent underabundance compared with the other ele-
ments in Figure 5. Na is an element that should have been enriched in the formation of a plagioclase-rich crust but that despite crustal enrichment is not even present in average solar system abundance levels.

A direct indication of the chemistry of the source regions of the lunar surface rocks is the initial $^{87}\text{Sr}/^{86}\text{Sr}$ ratio, i.e., the value of this ratio at the time of crystallization [Papanastassiou and Wasserburg, 1972a]. This quantity is very accurately determined and will vary depending on the contribution of $^{87}\text{Sr}$ from the radioactive decay of $^{87}\text{Rb}$ in the source regions of the rocks prior to their formation. In turn, this will depend on the Rb/Sr ratio of the source region. A high Rb/Sr in the source region will tend to give higher initial $^{87}\text{Sr}/^{86}\text{Sr}$ ratios and vice versa. As is shown in Figure 7, lunar rocks have initial $^{87}\text{Sr}/^{86}\text{Sr}$ ratios that are among the lowest known for any solar system material, even for lunar rocks that formed almost $1.5 \times 10^9$ yr after the solar system. The source regions are required to have Rb/Sr much less (~0.005) than average solar system values. The separation of Rb and Sr and presumably of all other volatile and refractory elements as well must have occurred very early in the history of the solar system, or else initial $^{87}\text{Sr}/^{86}\text{Sr}$ values much higher than those for meteorites would be observed.

Similarly, Pb isotopic data [Tera et al., 1974b] show that the source regions for lunar rocks are strongly depleted in Pb (a volatile element) compared to U and Th (refractory elements). But, more importantly, the Pb data also show that lunar rocks formed from parent materials which had Pb/U ratios much lower than those for the source regions of terrestrial rocks, conclusively showing a fundamental chemical difference between the earth and the moon.

**Volatile element abundance patterns.** As was discussed in the preceding section, elements such as Na, K, and Rb, which are only moderately volatile, appear to be depleted; thus it might be expected that elements that are highly volatile should be highly depleted. Figure 8, based primarily on data from Anders and his co-workers, shows that although volatile elements are depleted overall, there is a surprisingly large range of abundances for these elements in lunar rocks. The halogens...
A distinctive geochemical property is volatility. In addition to being volatile, Tl behaves as a lithophile element [Krahenbühl et al., 1973] and is known to correlate with alkalis in terrestrial igneous rocks [Jovenovic and Reed, 1974; Tatsumoto et al., 1973]. Similarly, Zn and Bi are chalcophilic, i.e., tend to concentrate in sulfide minerals. Except for F, which appears to behave somewhat independently for reasons that are not clear, these elements show a relatively restricted range of depletions of $10^{-4}$-$10^{-2}$ in mare basalts but show a wide range of concentrations (spanning 4 orders of magnitude for Tl) in highlands rocks. A large number of highlands rocks have 10-100 times higher concentrations than mare basalts. However, coarse-grained crystalline highlands rocks, such as the anorthosite rock 15415 or the plagioclase-olivine rock 76535, have extremely low abundances of volatile elements, as is illustrated specifically in Figure 8. The rocks (e.g., 66095) showing spectacular Tl enrichments (~1 on Figure 8), along with distinct enhancements of the other volatile elements (except F), are from the southern part of the Apollo 16 site; however, several different rock types are represented in this population [Krahenbühl et al., 1973; Morgan et al., 1974]. Rock 66095 is the famous ‘rusty rock’ [Taylor et al., 1973], which attracted attention because of prominent red stains due to the presence of hydrated ferric oxide. The rust and H$_2$O appear to be the result of oxidation of FeCl$_3$ in the rock by terrestrial oxygen and water vapor [Taylor et al., 1974; Epstein and Taylor, 1974]; nevertheless, the presence of a mineral like FeCl$_3$ (and ZnS [El Goresy et al., 1973]) in a lunar rock is of considerable interest. The Tl enrichments appear to be unrelated to the lithophile character of this element [Krahenbühl et al., 1973] and are probably too large to be ascribed to the addition of any known type of meteoritic material, although this may not be true for Bi. Moreover, 66095 contains excess Pb with a distinctly lunar isotopic composition [Nunes and Tatsumoto, 1973]. At least one of the Tl-rich rocks (60025) appears to contain indigenous lunar Xe, which is isotopically identical to the Xe in the terrestrial atmosphere and distinct from the Xe found in most meteorites or the solar wind [Lightner and Marti, 1974]. This is an additional argument for a lunar origin for the volatiles in the volatile-rich Apollo 16 rocks. Because these rocks are about $3.9 \times 10^9$ yr old (section B4), the volatiles must have been incorporated very early in lunar history. Thus these large volatile enrichments must be ascribed to outgassing of at least the outer portions of the moon during crystall formation [Krahenbühl et al., 1973], and rocks like 66095 are probably the closest lunar analogs to the terrestrial oceans and atmosphere. (The ancients had it backwards; the volatiles are in the highlands not in the maria!)

If outgassing has enhanced the abundances of volatile elements in lunar surface rocks, the fact that most rocks still have abundances that are $10^{-2}$-$10^{-1}$ of carbonaceous chondrites indicates either that the source material was highly depleted in volatile constituents or that loss mechanisms were very efficient in removing the outgassed volatiles from the moon. For example, ionization and acceleration of an atom by the solar wind can sweep material out of a tenuous lunar atmosphere [Manka and Michel, 1971]. This latter possibility may play some role, but the absence of any evidence for lunar H$_2$O, the overall low C content of lunar rocks, and the Pb isotopic data argue for the former alternative.

Volatile release was not confined to the highlands or to the early stages of lunar crustal formation, as is shown by the vesicularity of many mare basalts. However, the clearest indications of volatile release accompanying volcanism in the maria are the large enrichments in Zn, halogens, Pb [Tatsumoto et al., 1973; Silver, 1974], and other volatile elements in the Apollo 15 green glass (15426) and Apollo 17 orange glass (74220, Figure 8) samples. These are relatively homogeneous glasses, primarily in the form of 10- to 100-μm spherical shapes and have moderately well defined $^{40}$Ar ages [Huneke et al., 1973; Podosek and Huneke, 1973]. They are probably lunar volcanic glasses, and the associated volatile elements were incorporated at the time of eruption. There is evidence [Jovanovic and Reed, 1974; Tatsumoto et al., 1973; Silver, 1974; Grant et al., 1974] that the volatiles are on the surfaces of the spheres. However, the isotopic composition of the easily leached (Tatsumoto et al.) or volatilized (Silver) Pb in 74220 has a very high (nonradiogenic) $^{206}$Pb content. Thus the Pb

![Fig. 8. Volatile element abundance patterns in lunar rocks, expressed as the ratio of the measured concentration in lunar rocks to that of CI carbonaceous chondrites. Data for soils and soil breccias are not plotted except for the Apollo 17 orange glass (74220) and the Apollo 15 green glass (15426), which are regarded as igneous glasses and plotted along with mare basalts. Abundances are plotted relative to CI carbonaceous chondrites by using reference values from Ganapathy et al. [1970] and 120 ppm for F [Goldberg et al., 1975]. Data for F are from papers by Reed and his co-workers and by Wänke and his co-workers in the Lunar Science Conference proceedings plus work by Reed and Jovanovic [1973]. Similarly, data for other elements are from work by Anders and his co-workers and by Wänke and his co-workers in the Lunar Science Conference proceedings plus work by Reed and Jovanovic [1973]. Similarly, data for other elements are from work by Anders and his co-workers and by Wänke and his co-workers in the Lunar Science Conference proceedings plus work by Reed and Jovanovic [1973]. Similarly, data for other elements are from work by Anders and his co-workers and by Wänke and his co-workers in the Lunar Science Conference proceedings plus work by Reed and Jovanovic [1973]. Similarly, data for other elements are from work by Anders and his co-workers and by Wänke and his co-workers in the Lunar Science Conference proceedings plus work by Reed and Jovanovic [1973].
must have originated at the time of the orange glass eruption in a source region that had a much lower U/Pb ratio (i.e., a region much higher in volatile elements) than regions from which other lunar rocks were derived. Moreover, this source region must have remained isolated during the major period of crustal formation before $3.9 \times 10^9$ yr ago in order to preserve this low U/Pb. This is strong evidence for significant chemical heterogeneities in the lunar crust and/or mantle. It is interesting that as is true with the volatile-rich highlands rocks, the C and H$_2$O contents of the 74220 orange glass are extremely low [Epstein and Taylor, 1973]. This suggests that the moon formed from material with sufficient halogens, Zn, Tl, Bi, etc., that after being outgassed during crustal formation these elements can attain concentrations approaching carbonaceous chondrite levels in some rocks. However, corresponding C and H$_2$O enrichments are not observed, probably indicating that the initial lunar material was extremely depleted in C and H$_2$O.

**Siderophile element abundance patterns.** Lunar rocks are also depleted in siderophile elements, i.e., those elements that are known experimentally to concentrate in a metallic phase. Many of these elements (e.g., Ni and Ir) are highly nonvolatile; thus the overall depletion is undoubtedly related to their siderophile properties. Figure 9 summarizes the abundances of four key siderophile elements for which large bodies of data are available. Similar trends are present, as were noted for the volatile elements in the discussion of Figure 8. Although there is a wide spread in concentrations, the mare basalts show overall large depletions ($10^{-2}$-$10^{-4}$ of CI chondrites) despite the presence of small amounts ($\approx 0.1$%) of native metal in the rocks. Although there is also a wide spread, many highlands rocks show concentrations of these elements which are 1–2 orders of magnitude larger than those of mare basalts. There appears to be little discussion in the lunar science literature of the possibility that the enhanced highlands rock abundances are more typical of the moon as a whole and that the low abundances of mare rocks represent differentiation of a metallic phase. Most lunar chemists prefer an alternative interpretation, following Anders and his co-workers, in which the indigenous levels of siderophile elements are regarded as being low, comparable to those of the mare basalts. The larger abundances in highlands rocks are interpreted as residual ancient meteoritic material from the period of intense highlands cratering and in particular, from large basin-forming impacts. [Morgan et al., 1974, 1975]. Siderophile elements are comparatively abundant in many meteorites; thus even if the incoming projectile is completely volatilized, the siderophile element content of the residual material will still show the meteoritic contribution. The arguments for this interpretation are as follows:

1. There are some highlands rocks that have a siderophile element content as low as that found in mare basalts but are similar in bulk and lithophile element chemistry to the siderophile-rich highlands rocks [Morgan et al., 1975]. Many of the latter rocks are breccias, whereas the former are ‘crystal line’ (i.e., rocks showing an igneous texture, e.g., 76535) and pure anorthositic rocks, e.g., 15415. The siderophile-poor samples would represent uncontaminated highlands fragments that escaped brecciation during the period of intense bombardment. Crystalline rocks like 14310 or 68415, which have high siderophile contents, are regarded as rocks crystallized from an impact melt. The siderophile elements reside in metallic grains [Wlotzka et al., 1972; Ganapathy et al., 1973], which are a discrete component in the breccias and in this sense can be regarded as being extraneous to the silicate portions of the rock [Morgan et al., 1975].

2. Studies of radiation damage features indicate that a large number of highlands breccias, particularly those from Apollo 14, may contain small amounts of recognizable regolith materials, even though to a petrologist these may not be recognizable soil breccias [Hutchison et al., 1972; MacDougall et al., 1973]. Meteoritic contributions of siderophile elements to mare soils are well documented (see section C [Morgan et al., 1975; Baedecker et al., 1974]). The thermal metamorphism accompanying the formation of a breccia would destroy most of the normal indications of regolith materials (impact glasses, particle tracks, solar wind rare gases) but would not affect the meteoritic siderophile element contribution. Thus even if only a residual trace of regolith material is present, it is quite plausible that large enrichments in siderophile elements will also be found. It should be emphasized that these breccias were formed $3.9 \times 10^9$ yr ago; thus ancient regoliths are being discussed. A difficulty with the above argument is that a surprisingly large fraction of the highlands breccias would have to be composed of ancient regolith material.
3. Given the impact origin of highlands breccias, it is probably unreasonable to believe that all of the material originally in the impacting objects would be lost to the moon. A few percent by mass of the breccias is required to be from the projectile; this requirement is reasonable. From this point of view, siderophile enrichments in highlands breccias were to be expected, and it would have been more surprising not to have found them.

Although the above arguments are plausible, I do not think that they are sufficient to rule out completely an indigenous origin for siderophile elements in highlands rocks. In particular, it is striking that the rusty rock 66095 and the orange soil 74220, which are enriched in volatile elements (Figure 8) presumably of lunar origin, also have the high siderophile abundances (Figure 9), which are interpreted as being meteoritic. However, there are other rocks, e.g., 60025, that are rich in volatiles but very low in siderophile elements. Further, it is difficult to compile a list of arguments for an indigenous origin. In particular, it would be hard to understand the variability in the relative abundances of the various siderophile elements [Morgan et al., 1974, 1975]. This may require a variety of chemically distinct metallic source regions in the lunar crust, somewhat reminiscent of the 'plum pudding' theory of the origin of iron meteorites. Even when one allows for the possibility of inhomogeneous accretion of materials to form the moon, it is not clear that in a well-differentiated lunar crust such metal pockets could remain chemically isolated.

Regardless of the origin of the siderophile elements in highlands rocks, two very interesting general observations can be made:

1. Lunar surface rocks are depleted in siderophile elements by at least a factor of 10–100 (Figure 9) in comparison with average solar system material.

2. The relative abundances of the siderophiles [Morgan et al., 1974, 1975] among themselves are highly variable and do not match abundance patterns for contemporary meteorites.

The first observation undoubtedly indicates that a metal phase has been separated sometime in the early history of lunar surface material. A major question that remains is, exactly how did this happen, and where is the separated metal today? Perhaps siderophile elements are concentrated in a small lunar core (see the next section) or in widely disseminated pockets or plums in the deep lunar interior. However, given the limits on the mass of a metallic core set by the overall density and moment of inertia [Toksöz et al., 1974], unreasonable concentrations (percent levels) of siderophile elements in the core would be required for the moon as a whole to have average solar system abundances of these elements. It is more generally accepted that the separation occurred in the solar nebula [Anders et al., 1971] and that the residual metal is nowhere near the orbit of the earth. It probably cannot be totally ruled out that the missing siderophiles reside today in the earth’s core, although siderophile abundance patterns in terrestrial basalts appear to be distinct from those in lunar mare basalts [Anders et al., 1971].

Observation 2 above is of great importance because the relative abundances of the lithophile elements are qualitatively similar to those found in achondritic meteorites. However, the lunar siderophile abundance patterns suggest that there were early solar system materials that are chemically very different from those discovered through meteorite studies and that many fundamentally different materials remain to be discovered in the solar system.

**Summary of lunar chemistry.** In summary, because essentially total escape of volatile constituents from the moon during crustal formation appears to be less plausible, the underabundance of volatile elements compared with either lithophile refractory elements or carbonaceous chondrites is usually regarded as a property of the moon as a whole or at least the regions that were differentiated to form the lunar crust. Similarly, the low abundances of siderophile elements are best regarded as reflecting the separation of a metal phase from lunar material prior to the formation of the moon. This is undoubtedly a key factor in explaining the low density of the moon. The implied high-temperature history of lunar material can readily be reconciled with current ideas concerning chemical processes in the solar nebula, and several models have been discussed [Ganapathy and Anders, 1974; Anderson, 1973a; Wänke et al., 1974; Grossman et al., 1974; Taylor and Jakes, 1974].

3. **Density and Internal Structure of the Moon**

There are several excellent recent reviews of these subjects [Toksöz et al., 1974; Nakamura et al., 1974a; Lammlein et al., 1974]; they are also summarized here because of their importance. There are three physical properties of the moon that are known as a function of depth from direct measurement: the P and S wave seismic velocities [Toksöz et al., 1974; Nakamura et al., 1974a] and the electrical conductivity [Dyal et al., 1974]. The conductivity profile can be used to estimate the temperature profile for an assumed lunar interior composition. As for the earth, seismology provides the best look at the interior structure of the moon. Three types of seismic signals have been utilized: (1) man-made impacts of spacecraft [Toksöz et al., 1974], (2) meteorite impacts [Latham et al., 1973], and (3) lunar seismic events [Lammlein et al., 1974]. Most moonquakes are very weak by terrestrial standards (typically Richter magnitude 1) and are much less frequent (the seismic energy release of the moon is estimated to be $10^{-14}$ that of the earth). However, rare 'high-frequency teleseismic' events have reached magnitude 4 [Nakamura et al., 1974b].

The compressional wave seismic velocity profile in the upper 100 km of the moon (Figure 10), determined primarily from spacecraft impact data, shows two discontinuities, those at 20 and 55 km. There is good evidence for both discontinuities, and the case for the 55-km discontinuity is particularly strong [Toksöz et al., 1974]. Below 60 km it is not possible to fit all of the observed travel time data with a single velocity. Two choices are possible: 7.7 or 9.0 km/s. If the higher velocity is real, it either must be a regional layer or must be very thin (~40 km) in order to match the velocity at deeper regions in the moon (see discussion below). The 55-km discontinuity is taken to be the base of the lunar crust and is probably the lunar counterpart of the terrestrial Mohoroviči? discontinuity. The particular profile shown in Figure 10 applies to the western mare regions and may not be typical of the whole moon. The 20-km discontinuity can be interpreted as the depth of the mare basalts in this region. Laboratory P wave velocity measurements on lunar samples show no clear distinction between various lunar rock types (see compilation by Toksöz et al. [1974]); thus it is not possible to make firm compositional assignments to the observed crustal layers. However, the laboratory velocities are in approximate agreement with the measured crustal velocities, although the returned samples require higher pressures (equivalent to ~50-km depths) to reach the observed crustal velocity in the 20- to 50-km depth range (see also the paper by Mizutani and Osako [1974]). This is probably because returned lunar samples are more highly frac-
Fig. 10. Compressional wave velocity profile in the upper crust from Toksöz et al. [1974]. The velocities drop fairly rapidly above 20 km owing to increasing amounts of fracturing (impact?) of the crustal rocks. The discontinuity at 20 km marks the depth of the mare basalts, whereas the discontinuity at 55 km represents the base of the crust. There are two possible values for the velocity at the top of the mantle, as is indicated.

tured than the actual lunar crustal rocks, requiring higher pressure for pore closure.

However, the densities of lunar crustal rocks provide some constraints on the composition of the deep crust and on the nature of the 55-km discontinuity. The densities of mare basalts are relatively high, 3.3–3.4 g/cm³, reflecting the high Fe and Ti contents. Thus mare basalts are surprising materials to find on a planet with average density of 3.34 g/cm³. However, the feldspar-rich highlands rocks will have densities of 2.8–3.1 g/cm³ and are therefore much more plausible crustal materials. All crustal rocks are feldspar-bearing and are thus unstable with respect to phase transitions at deep interior pressures [Ringwood and Essene, 1970; Anderson, 1973a, b]. However, prediction of the exact depths at which these transitions should occur is complicated by uncertainties in the temperature gradient and by the fact that there is a wide pressure range for the transition for feldspar-rich rocks [Anderson, 1973a]. Nevertheless, it is certain that mare basalts will undergo phase changes to rocks with densities of 3.5–3.7 g/cm³ [Ringwood and Essene, 1970; Green et al., 1971], which are incompatible with the overall lunar density. Thus mare basalts cannot make up the deep interior regions of the moon. The situation for feldspar-rich highlands rocks is more complicated in that it may be possible for Ca- and Al-rich rocks of acceptable density to exist at depths of at least several hundred kilometers [Anderson, 1973a, b]. An interesting possibility is that the 9.0-km/s 'high-velocity zone' might represent a phase change from a feldspar-rich to a garnet-rich zone at 60 km but then revert back to a feldspar-bearing assemblage at a slightly greater depth because of the increase in temperature [Toksöz et al., 1974].

Figure 11 shows the seismic velocity structure of the deep interior of the moon, based on meteorite impacts and internal lunar seismic events. Velocity profiles calculated by Toksöz et al. [1974] and Nakamura et al. [1974a] are in overall good agreement, although the two groups differ in that Nakamura et al. report a discontinuity at 300 km that they propose as the base of the zone from which the crust has been differentiated. Magnesium-rich olivine and orthopyroxene mixtures provide a good match for the average seismic velocity and the density of the mantle above ~700 km from the center. Pure olivine compositions are possible but not pure orthopyroxene. The ~8-km/s average mantle P velocity is somewhat low in comparison with estimates for Ca- and Al-rich rocks [Anderson and Kovach, 1972] unless temperatures higher than those estimated by Toksöz et al. [1974] are adopted.

For seismic ray paths that pass within about 700 km of the center, shear waves are highly attenuated (Figure 11). The effect on the P wave velocity is much less. This behavior is similar to that of the asthenosphere (low-velocity zone) of the earth and may also represent a partially molten zone in the lunar interior. Alternatively, it could represent a volatile-rich, undifferentiated lunar 'core.' This latter alternative, if it were true, would require a spectacular chemical contrast between the inner and the outer portions of the moon and would probably indicate some kind of inhomogeneous accretion for the moon. On the other hand, a partially molten lunar interior

![Figure 11. Velocity profiles in the deep lunar interior as determined by natural seismic events. Profiles as calculated by Toksöz et al. and by Nakamura et al. are in reasonable agreement. The strong attenuation of S waves at 7- to 800-km radius is interpreted as the onset of partial melting. The dip in P velocity at 2- to 300-km radius may indicate a small molten core. Figure is from Toksöz et al. [1974].](image-url)
does not appear to violate any geophysical or geochemical constant and can quite easily be reconciled with thermal history calculations [Tokès et al., 1974]. This is the more plausible and more widely accepted alternative at present.

A single meteorite impact event on the extreme back side of the moon has shown a decrease in $P$ wave velocities for rays traveling through the innermost 20% of the moon (Figure 11) [Nakamura et al., 1974a]. No $S$ waves were detected. If additional events confirm the drop in $P$ wave velocity, it will indicate that the moon does have a small liquid core. A molten iron core of up to 30% of the lunar radius (5% by mass) is compatible with the lunar density and moment of inertia. Similarly, an FeS core of 40% in radius and 8% by mass would be possible [Tokès et al., 1974]. Thus the core indicated by the Nakamura et al. data could be either metal or sulfide [Brett, 1973]. Conceivably, this is the home of some of the siderophile and/or chalcophile elements that are highly depleted in lunar surface rocks.

The longevity of the lunar seismic network has paid off handsomely.

4. Evolution of the Moon

Figure 7 displays the age distribution of lunar rocks as obtained from $^\text{40}Rb-\text{Sr}$ internal isochrons [Tera et al., 1974a]. This remarkable age pattern was totally unexpected and warrants detailed discussion. (All ages in the following discussion will be given in units of $10^9$ yr.)

At least one lunar rock (72417) has survived the initial crustal formation. This is a dunite (95% olivine) that was sampled as a fragment from a breccia boulder at the base of the highlands massifs during Apollo 17. It has an Rb-Sr isochron age of $4.6 \times 10^9$ yr, although it is shocked [Albee et al., 1974]. Petrologically, it appears to be a cumulate rock that formed in at least the deeper parts of the crust. The more recent violent history of the region in which this rock formed failed to disturb completely the initial Rb-Sr systematics. This rock in itself demonstrates very early igneous activity on the moon, probably marking the initial differentiation of the lunar crust in the period 4.4-4.6. Other rocks, e.g., 12013 [Lunatic Asylum, 1970], 65015 [Papanastassiou and Wasserburg, 1972b], and 76055 [Tera et al., 1974b], can be interpreted as rocks originally formed at 4.4-4.6 but extensively reheated at about 3.9. Only a few rocks have ages in the time span between 4.0 and 4.6 [Bogard et al., 1974; Husain and Schaeffer, 1975]. The record of this period of lunar history appears for the most part to have been totally obliterated. The intense activity terminated in the period 3.85-4.0 on the basis of ages of highlands rocks from Apollo 14, 15, 16, and 17 and Luna 20, and it appears that most highlands rocks have ages of 3.85-4.0, although more samples are required to verify this, particularly from the lunar far side.

Fra Mauro (Apollo 14) rocks are samples of a region interpreted as a ridge of ejecta produced by the impact event that formed the Imbrium basin (Figure 1). This interpretation is vital to our present understanding of the moon and should be subjected to further photogeologic study. As judged from the 3.2 ages (Figure 7) for Apollo 15 mare basalts, the lava flows that filled the basin to produce Mare Imbrium occurred long after the formation of the Fra Mauro rocks. Strictly speaking, the age of the Imbrium impact is bracketed between 3.2 and 3.85, but 3.85 is usually adopted as the time of the Imbrium impact because (1) an impact large enough to produce a 500-km-diameter crater should produce strong heating and some melting of the ejecta, (2) most of the Apollo 14 rocks are impact breccias, and (3) K-Ar ages, which are sensitive to reheating, are also in the range 3.85-3.95 [Turner, 1971]. Mare Imbrium is the second youngest (after Mare Orientale) of approximately 40 major lunar impact basins [Howard et al., 1974]. The intense bombardment of the moon by these and innumerable smaller impacts sculptured the lunar surface in the period between 4.6 and 3.9, producing the rugged highlands topography and obliterating most of the record of lunar igneous activity in that period. The ages of mare basalts are only slightly younger than those of highlands rocks; this finding is very surprising in view of the very large difference in crater densities. Quantitative comparison of the ages (Figure 7) for the various landing sites and the corresponding density of large impact craters in the same regions [Boyle et al., 1974] shows that there is a large increase in crater density for times older than 3.5. It had long been acknowledged that impact rates might be higher earlier in the history of the solar system, but such a dramatic increase was totally unexpected. Further, because the collisional lifetimes of objects in earth-crossing orbits are estimated to be $10^9-10^{10}$ yr at most [Wetherill, 1974], it was also a major surprise to learn that the intense early bombardment of the moon (and the earth), as recorded in the lunar highlands crater density, had extended to times up to $0.7 \times 10^9$ yr after the formation of the solar system. It may be that the moon was subjected to an intense pulse of bombardment at $4 \times 10^8$ yr ('terminal cataclysm' [Tera et al., 1974b; G. W. Wetherill, unpublished manuscript, 1974]).

Basalt samples from five of the major maria (Figure 7) show a relatively small range in age from 3.3 (Apollo 12 and 15) to $3.9 \times 10^9$ yr (Apollo 11 low K rocks [Turner, 1970; Eberhardt et al., 1974]). The major point is that there appears to be a cessation of igneous activity as $\sim 3.2$ [Papanastassiou and Wasserburg, 1971]. Comparison of relative crater densities for the youngest regions on the moon with those of the mare landing sites, when one allows for a time-dependent bombarding flux, indicate that the youngest mare flows would be $\sim 2 \times 10^8$ yr [Boyle et al., 1974].

A unique, unanticipated thermal history for the moon is indicated. Independent of specific models, the differentiation accompanying crustal formation must have begun no later than $\sim 4.4$. Whether crustal formation occurred suddenly at 4.4-4.5 or continuously down to 4.0 is not clear at present. There are some rocks intermediate in age, particularly the troctolite 76535, which appears to be a deep crustal cumulate rock that formed at about 4.3 [Bogard et al., 1974; Husain and Schaeffer, 1975], but it is not clear whether or not these are rare interlopers.

There is no evidence for highlands magmas of either internal or impact origin at times later than 3.85 (determined by 68415 [Papanastassiou and Wasserburg, 1972b]). The time of initial lava flows in the mare basins is unknown, but the oldest sampled surface flows are $\sim 3.9$. Basin filling appears to have terminated at about 2-3. And then nothing? Today the moon is extremely quiet. It is solid down to depths of 1000 km with seismic activity $10^{-14}$ that of the earth (see the previous section). Why should (and how can) a planetary-sized body go through $1.5 \times 10^9$ yr of intense igneous activity and then stop? The earth has been thermally violent throughout its history and continues active until the present day. The parent bodies of most meteorites formed very rapidly $4.6 \times 10^9$ yr ago and showed no further activity. A specific lunar thermal history that can reproduce these general requirements is a matter of theoretical calculation, and many papers on this subject have been written [e.g., Tokès and Solomon, 1973], but there
is a general consensus among the various calculations that the initial temperature gradient in the moon was peaked toward the surface, permitting early igneous activity, but that the inside was relatively cool, permitting the deeper regions to cool fairly rapidly and igneous activity to cease.

5. Summary and Epilog

The preceding discussion attempted to outline what progress has been made in answering the list of 'fundamental problems' given in the introduction. In addition to summarizing this discussion it is also important to consider what remains to be done.

We now have a good basic characterization of the nature of highlands and mare materials. Thus it is not premature to regard these as the data that must be described by petrogenetic models, and inspection of the Proceedings of the Fifth Lunar Science Conference reveals that much work has already been initiated. Key problems are (1) the nature of the source regions of the mare basalts and the role of near-surface differentiation [Hays and Walker, 1975], (2) the possible association of the highlands rocks with major impact basins, and (3) the origin of the highlands crystalline rocks, allowing for the possibility of impact melting. However, because of the pace of the lunar missions it was not possible to distribute all lunar samples for study; thus it is an equally important post-Apollo task to ensure that fundamentally different types of lunar materials have not been overlooked and that the remaining samples are in accord with the patterns established to date.

The evidence is very strong that the moon is a differentiated planet, as is the earth. The moon has a crust enriched in lithophile elements that was derived from a source region depleted in volatile substances, including the alkali elements. Key element ratios, e.g., K/U, appear to be approximately preserved in planetary differentiation processes and thus can be used as 'planetary constants.' Interestingly, some of these ratios, e.g., K/U and Pb/U, are different for the moon and the earth; this puts a serious constraint on the relative origins of the two planets. The experiments of Metzger et al. [1974] have demonstrated that by using the gamma rays from their radioactive decay, K, U, and Th can be determined from an orbiting spacecraft, documenting an important chemical probe for planets in general. In the western mare regions the lunar crust extends to a depth of ~60 km, underlying 20 km of mare basalt [Toksöz et al., 1974]. The lunar crust is much thicker than that of the earth and is probably composed primarily of feldspar-rich rocks. The composition of the lunar interior cannot be uniquely determined but is strongly constrained by the seismic velocity and density. For example, mixtures of Mg-rich olivine and pyroxene match the properties of the lunar interior well, but a fairly narrow range of composition is required. The amount of the lunar interior that must be differentiated to produce the crust is model dependent. In general, estimates could range from zero (for an extreme inhomogeneous accretion model defended by no one at present) to the whole moon [Wood, 1975 (this is compatible with thermal models); Toksöz and Solomon, 1973]. More typical estimates tend to be in the range of hundreds of kilometers, e.g., 400 km [Ringwood and Green, 1974]. A seismic velocity discontinuity has been proposed at 300-km depth by Nakamura et al. [1974a], which might mark the depth of the zone from which the crust was differentiated.

The strong attenuation of S waves beneath 1000-km depth might indicate a fundamentally different, possibly volatile-rich interior; however, a partially molten zone, analogous to the earth's asthenosphere, is the more popular interpretation. Some evidence exists [Nakamura et al., 1974a] for a small molten core that could be either Fe or FeS [Toksöz et al., 1974]. The presence of a lunar core may account for some of the depletions of siderophile and/or chalcophile elements in lunar surface rocks.

The prime future task is to convert the qualitative observations cited here into a quantitative chemical history for the moon. Many ideas on this subject have already been proposed (see, for example, volume 2 of the Proceedings of the Fifth Lunar Science Conference). Refined seismic data on the nature of the lunar interior are required, and the permanent existence of a seismic network is crucial.

The evolutionary history of the moon was characterized by a period of relatively intense igneous activity that terminated 2-3 \times 10^9 yr ago. Intense meteoritic bombardment of the lunar surface has obscured the igneous record for times older than about 4 \times 10^9 yr. Most thermal history calculations for the moon [Toksöz and Solomon, 1973] seem to require an initial temperature gradient peaked toward the surface. It is very important to see whether this type of temperature gradient is inescapable in order to account for the present structure, composition, and evolution of the moon. Two methods have been proposed to account for such a gradient: (1) rapid accretion of small, cold grains in which the rate of gravitational energy release in the terminal stages of growth is too rapid for efficient radiative cooling [Hanks and Anderson, 1968] or (2) inductive heating of an initially cold moon in an intense early solar wind [Sonett et al., 1969]. Both of these mechanisms have the common property that the moon formed basically cold and was heated only in the final stages of growth (rapid accretion) or after formation (inductive heating). Contemporary planetary scientists have generally agreed that some type of cold accretion is the most plausible way for planets to form, but if it can be shown that this must be true for the moon, this will constitute a major advance.

C. UNANTICIPATED RESULTS FROM APOLLO

The results to be presented in this section are equally as important as the topics discussed in section B. From the list given in the introduction, items 1 and 2 have already been discussed in sections B4 and B1, respectively. The remaining items will be discussed below in their order in the original list.

Ancient Paleomagnetic Field

Lunar magnetism is covered in recent detailed reviews by Fuller [1974] and Dyal et al. [1974a]. Based on measurements from the orbiting subsatellite launched by Apollo 15, the present-day global dipole field of the moon is less than 10^-4 of that of the earth [Russell et al., 1974]. However, local surface fields approximately 1% of those of the earth were found during Apollo 16 [Dyal et al., 1973], and regions of high magnetic fields have been observed from orbit, particularly near the crater Van de Graaff on the lunar far side [Sharp et al., 1973]. These high local fields are a reflection of a high degree of natural remanent magnetism observed in many lunar samples. The remanent magnetism is usually interpreted as being thermal in origin, thus indicating the presence of an ambient magnetic field at the time that the lunar samples cooled below the Curie point (750°-800°C). Paleointensity estimates of the required fields give values of 0.05 to several times the terrestrial field [Gose et al., 1973; Stephenson et al., 1974]. Some of the measurements are on breccias (e.g., 15498) whose most recent time of heating is not clear and could be quite recent.
Other measurements are on Apollo 16 crystalline rocks and mare basalt samples [Stephenson et al., 1974] for which it is quite likely that the time of final cooling is given by the crystallization age. Taken at face value, these data qualitatively indicate that sizable magnetic fields existed at the lunar surface 3-4 × 10^9 yr ago, which are totally absent today. However, quantitative estimates of paleofields are complicated by the effects of the detailed thermal history of a given rock, in particular, shock [Fuller, 1974; Cisowski et al., 1974; Brecher et al., 1974]. For example, Apollo 17 mare basalts of the same age show an order of magnitude variation in the derived paleomagnetic intensities even from the same laboratory [Stephenson et al., 1974; Brecher et al., 1974].

Depending on how it is assumed to originate, an ancient lunar magnetic field can lead to very different views of the early thermal history of the moon. If the field were due to a molten metal core [Pearce et al., 1972], then the interior of the moon would initially have been hot (1600°-1700°C), and it is not clear whether the outer portions would cool rapidly enough to cause igneous activity to cease at ~3 × 10^9 yr ago and to support the gravitational anomalies represented by the mascons [Toksöz and Solomon, 1973; Toksöz and Johnston, 1975]. Alternatively, it has been postulated that the moon was initially below the Curie point at all portions of the interior and was magnetized by a transient early solar system magnetic field [Strangway and Sharp, 1974]. The moon was presumably still magnetized in the interior throughout the duration of mare filling, but as the interior temperature rose through the Curie point, the magnetic field vanished. Resolution of these alternatives appears to require reliable quantitative paleomagnetic determinations in order to determine the time dependence of the paleofield.

If there is a small molten core in the inner 100-200 km of the moon today (section B3), it does not appear to generate an appreciable magnetic field. According to Brett [1973] an Fe-FeS eutectic mixture has sufficient conductivity and magnetic permeability to produce a magnetic field for the present rotation rate of the moon. A much larger rotation rate would be required for a pure FeS core. If the ancient lunar magnetic field were due to a molten core and if the core still exists, the moon is required to have had a much more rapid rate of rotation in the past. The decay of the ancient field might reflect a decrease in the rotation rate.

Exotic Components in the Regolith

With the possible exception of a few Apollo 15 rocks from the edge of Hadley Rille, which may be bedrock samples, all returned lunar samples were impact-derived fragments sampled from the debris layer (regolith) that makes up the outer ~10 m of the lunar surface. Except for impact-produced glasses the finer-sized (~1 mm) particles (soils) might be expected to be no more than comminuted material from the larger rock fragments. To a first approximation this is true; however, the fascinating area of 'regolith studies' derives much of its energy from the fact that a lunar soil contains much more than broken rock fragments. Only a few examples of research from this area are discussed.

Kreep. The first lunar soil from Apollo 11, 10084, had Rb and Sr contents and a Sr isotopic composition that could not be explained by mixtures of the local rocks [Papanastassiou et al., 1970]. An 'exotic component' that was comparatively rich in Rb was required. Relatively small, but definite enrichments in K, Cs, and Ba were also required [Tera et al., 1970]. Studies of millimeter-sized fragments revealed rare alkali field-spar-glass-low Ca pyroxene fragments [Albee and Chodos, 1970], which could account for the chemical and isotopic properties in the soil. These fragments were the first evidence for highly differentiated, graniticlike materials in the lunar crust. Similar studies of Apollo 11 coarse fines led to the discovery of anorthositic fragments, which are probably ejecta from highlands areas into the maria [Wood et al., 1970]. Apollo 12 returned soils for which the presence of an exotic component was no longer subtle. The K contents of Apollo 12 soils are 3-5 times higher than those of the local mare basalts owing to the presence of light-colored glassy particles named Kreep, because they were rich in many lithophile elements, including K, rare earths, and phosphorous [Meyer et al., 1971]. A single large rock, 12013, which is the closest lunar analog to a terrestrial granite, was also returned [Lunatic Asylum, 1970; Drake et al., 1970]. The Apollo 12 site was located on a light-colored ray from the large crater Copernicus, 370 km distant, and this is a plausible source for these materials. These studies emphasize the importance of lateral transport by impact for the moon and presumably for other atmosphereless planets. Because of lateral transport, a sample from any one spot gives information about materials from a much wider region.

Missions subsequent to Apollo 12 returned many K-rich rocks that have been called Kreep, although it is not clear that they represent a single rock type. In missions where there were local sources of K-rich rocks (Apollo 14, 16, and 17), it has been possible to account for the lithophile element chemistry of the soils in terms of mixtures of local rocks [Warner et al., 1974; Rhodes et al., 1974]. For the mare soils from Apollo 15 an external component is again required. Orbital gamma ray data show that the extremely K-, U-, and Th-rich areas tend to be concentrated in the near side regions around Mare Imbrium [Metzger et al., 1974].

Implanted solar ions and regolith mixing. Highly volatile elements, such as rare gases, are essentially absent from lunar rocks but are relatively abundant constituents in the solar wind. In the absence of an atmosphere or a significant global magnetic field the solar wind impinges directly on the lunar surface. Given their low energy (~1 keV/nucleon), the solar wind ions will penetrate only a few hundred angstroms and thus will not be present in interior rock samples. However, because of mixing of the regolith by small impacts, many of the grains of a soil sample will have been exposed directly to the solar wind. Consequently, the concentrations of rare gases in soil samples are many orders of magnitude higher than those in rocks [Eberhard et al., 1972]. Moreover, the isotopic compositions of the rare gases from interior rock samples show that they are formed primarily from galactic cosmic ray nuclear reactions, and these isotopic compositions are very distinct from those found in soils. Thus the solar wind rare gases are qualitative tracers for exposure of material to the outermost layers of the lunar surface. The efficiency of the impact mixing process is shown by the fact that roughly uniform solar wind gas contents are found at depths down to 3 m in the deep core samples, indicating that regardless of present depth, most of the material at one time resided on the lunar surface [Bogard et al., 1973; Eberhardt et al., 1974]. Similar conclusions are reached from measurements of tracks produced by solar flare heavy ions (10- to 100-MeV/nucleon Fe nuclei) that penetrate only ~100 μm, except that evidence of discrete depositional events is found in some cases [Arthennius et al., 1971; Fleischer et al., 1971; Crozaz et al., 1974]. The best information on the depositional history of the core samples is
provided by studies of neutron capture products because neutron capture occurs primarily at meter depths rather than at the surface [Russ et al., 1972; Russ, 1973; Curtis and Wasserburg, 1975] and because neutron capture rates are well known [Woolum et al., 1975].

The relative abundances of the rare gases show considerable variation among lunar soils and also for mineral separates of a given soil [Hintenberger et al., 1974]. Moreover, the lighter rare gases appear to be depleted in comparison with either estimated solar system abundances or the relative He/Ne/Ar abundances measured directly in the solar wind foil experiments during Apollo 11–16 [Geiss, 1973]. This shows that at least the lighter rare gases are not quantitatively retained. The order of magnitude of the solar wind Xe concentrations is consistent with no loss of solar wind Xe from the moon over \( 4 \times 10^9 \) yr, if the Xe is assumed to be distributed through a typical regolith depth of \( 4 \) m [Burnett et al., 1971a; Geiss, 1973]; however, by noting that the amounts of solar wind Xe are high in relation to the amounts of Xe produced by cosmic ray nuclear reactions, Geiss [1973] has proposed an increase in the solar wind flux in the past. Other authors have emphasized that the concentrations of cosmic ray reaction products, particularly those due to neutron capture, are low in comparison with what would be predicted from simple ideas of regolith mixing [Eugster et al., 1970; Burnett and Woolum, 1974; Curtis and Wasserburg, 1975]. This may or may not invalidate the conclusion of Geiss depending on the origin of the low concentrations of cosmic ray reaction products. More sophisticated estimates of regolith vertical mixing rates [Gault et al., 1974] appear too slow to explain the observed cosmic ray product data, and this is an area of intensive research at present.

Large concentrations of H (approaching \( 1 \) cm\(^3\) H\(_2\)/g) are found in lunar soils, undoubtedly of solar wind origin [Epstein and Taylor, 1973]. Most interestingly, the hydrogen appears to be D free (less than 0.1 of the terrestrial D/H ratio), confirming theoretical astrophysical predictions that D would be destroyed by thermonuclear reactions in the early stages of solar evolution. Since the solar D is converted to \(^4\)He, the solar wind \(^3\)He/\(^4\)He ratio (as measured in the solar wind foils), combined with an estimate of the solar \(^3\)He/\(^4\)He ratio, gives an upper limit for the D/H ratio of \( 3 \times 10^{-8} \), which is 5 times less than the terrestrial ratio, showing that the formation of the earth resulted in extensive isotopic fractionation of H [Geiss, 1973].

Both the absolute and the relative C and N concentrations in lunar soils are consistent with the bulk of these elements also being of wind origin [Kothari and Goel, 1973]. This is actually somewhat surprising, and the lack of a clear lunar component indicates, as was discussed in section B2, that lunar crustal enrichment in other volatile elements, such as Zn and Tl, was not accompanied by N, C, and H\(_2\)O enrichment, showing that the moon as a whole is very depleted in these. Also, the lack of a clear meteoritic contribution is important, as is discussed in the following section.

The chemistry of C in a lunar soil sample is complicated but fascinating. Most of the carbon released in a pyrolysis experiment is emitted as CO, but this may be formed during the heating [DesMarais et al., 1973]. Some CH\(_4\) is present, because this gas is released when lunar soils are attacked with deuterated acids that contain no hydrogen [Cadogan et al., 1972]. The presence of 'carbide'-like material is inferred because CD\(_2\) is also released in these experiments. The actual nature of the C compounds formed by solar wind implantation may be clarified by simulation experiments now underway in many laboratories.

**Meteoritic material.** For all the mare sites it has been found that the concentrations of volatile and siderophile elements in soils are much higher, by factors of 10–100, than those found in the mare basalts [Morgan et al., 1975; Baecker et al., 1974]. Because of high concentrations of many of these elements in highlands rocks it is more difficult to establish corresponding enrichments in Apollo 14 and 16 soils. Solar wind contributions should be negligible; thus there are three plausible sources for the excesses: (1) lateral transport from highlands regions where rocks rich in these elements are found, (2) local volcanic activity, and (3) residual meteoritic material. All three of these sources are viable mechanisms for the volatile elements and may contribute to some degree. For siderophile elements there is no evidence that under lunar conditions they would be concentrated in volcanic vapors or pyroclastic eruptions, and so alternative 2 looks less probable. Similarly, the mare soil concentrations are comparable (\( \sim 10^{-4} \) carbonaceous chondrite levels) to the volatile-rich highlands rocks. Unless much of the mare soil is really of highlands origin, lateral transport cannot explain the siderophile element concentrations. This same argument may not hold for volatile elements, because it is conceivable that these can be 'distilled' from the highlands rocks during impact, and once they are in the lunar atmosphere, they may be deposited and 'released' from surfaces several times before becoming chemically 'fixed.' This would cause volatile elements to be dispersed more widely following impact volatilization than less volatile constituents. Thus for siderophile elements most lunar scientists accept a meteoritic origin, at least for mare soils. This is further supported by the relative abundances of the siderophile elements, which as is shown in Figure 12, correspond to those found in CI carbonaceous chondrites and are distinct from those found in highlands rock samples. Approximately 1% of meteoritic material in a mare soil is indicated. Many volatile elements, e.g., Sb, Se, Ag, Te, and Bi, despite the possibility of alternative sources, show abundances of \( \sim 10^{-4} \) CI levels, also compatible with being primarily of meteoritic origin [Morgan et al., 1975]. There is a general scatter of about a factor of 2 in the CI-normalized abundances, with no consistent indication of systematic variations of siderophile relative to volatile elements. If 1% of CI meteoritic material were added to mare soils, the resulting C and N concentrations should be 300 and 20 ppm, respectively. This presents no problem for N, but typical mare soil carbon contents are 50–200 ppm [Moore et al., 1974]; moreover, the carbon abundances can be explained by solar wind contributions. It appears that either (1) the added meteoritic material is not predominantly CI carbonaceous chondrites but another type of meteorite with less C but with similar relative abundances of siderophile elements [Morgan et al., 1975], (2) C is selectively lost from the moon during impact, in comparison with other volatile elements, (3) C is preferentially lost from the regolith after deposition by diffusion or solar wind sputtering, or (4) we are being fooled by the consistency in relative abundances (Figure 12), and all of the siderophile and volatile elements in the soils are lunar in origin from an as yet unforeseen source. The consensus among workers in this area is that explanation 2 is correct, but all of these alternatives and the general question of the sources and chemical distribution of C and N in lunar soils have important implications and deserve further study. Item 4 is perhaps the most exotic because it would imply that there is essentially no
record whatsoever left on the moon of all the meteorites that produced the intensely cratered terrain.

**Fundamentally Different Material on Lunar Surface Layers—Properties of the Outer Micron**

The preceding section indicates that many of the interesting properties of lunar soils are related to the possibility of deposits on the surfaces of grains; thus the present topic really cannot cleanly be separated from the general area of regolith studies. I have arbitrarily singled out the physical and chemical properties of the first micron of the lunar surface as viewed from space for special discussion, because I feel that it is a very interesting area for further research.

The outer micron of the lunar surface is subjected to a bewildering variety of processes: (1) solar wind implantation, (2) deposition of atoms from a transient lunar atmosphere produced by impact volatilization or outgassing events, (3) ‘reimplantation’ of ions produced by ionization of atmospheric constituents, (4) thermal diffusion of the emplaced species (probably radiation damage controlled), (5) chemical changes (reduction) during or subsequent to emplacement, (6) deposition of submicron, molten ‘accretionary’ particles, (7) erosion of the surfaces by solar wind sputtering, and (8) destruction of the surface layers by impact melting with redistribution of surface constituents throughout the volume of the particles (glasses and agglutinates) formed.

Surfaces exposed to the solar wind are expected to become saturated \((10^{14} - 10^{15} \text{ ions/cm}^2)\) with implanted H and He in times of 10–1000 yr [Bibring et al., 1974; Maurette and Price, 1975]. Thus the chemistry of the outer micron should be heavily influenced by implanted solar wind. Further, as first documented by the studies of Borg et al. [1971], the implantation process systematically destroys crystal lattices, producing an amorphous coating with thickness comparable to the solar wind implantation depths, 200–500 Å. These authors showed that a surprisingly large number of \(\sim 1\)-μm soil grains had amorphous coatings when they were viewed under a high-voltage transmission electron microscope. The amorphous layers and the constituents that they contain have a finite lifetime of \(\sim 10^5\) yr (while directly exposed on the surface) because the production of the amorphous layer by implantation and destruction by sputtering are in equilibrium. Once it is buried, even a few microns, the lifetime of the layer itself is determined by its recrystallization rate. The concentrations and distribution of the constituents are determined by diffusion and chemical reaction rates. Little is known about any of these rates. Preferential loss, presumably by diffusion, of light constituents is shown by the low measured H/He and He/Xe ratios, which are about 1% of those estimated from solar abundances [Geiss, 1973]. As far back as Apollo 11, evidence for postimplantation redistribution of solar wind rare gases was shown by etching experiments [Eberhardt et al., 1970] and by mass spectrometric measurement of rare gases released during electron beam volatilization of individual soil grains [Kirsten et al., 1970]. Both experiments showed that appreciable concentrations of rare gases existed at depths up to 1 μm, or roughly 100 times the solar wind penetration depth. Figure 13 shows a quantitative profile of redistributed solar wind H, as measured using a nuclear resonant reaction technique for a surface sample of lunar rock [Leich et al., 1974]. Relatively large H concentrations are observed to depths as great as 0.4 μm, or over 10 times the range of solar wind protons. Despite the apparent redistribution the concentrations are still large in comparison with what would be expected from estimates of H diffusion coefficients in silicates, suggesting that diffusion rates are much lower in the heavily radiation-damaged regions and that the depth profile at depths greater than 500 Å is a reflection of the density of radiation-produced ‘traps’ in the mineral lattices. In the amorphous layer above 500 Å, diffusion is rapid again, and the H concentration drops rapidly. It is quite clear that the physical and chemical properties of the outer micron are dominated by radiation damage effects.

The best evidence for surface deposits is the observation of S-rich coatings by Auger spectroscopy on Apollo 17 orange glass spheres [Grant et al., 1974]. This was a favorable case because the orange glass has had little if any solar wind ex-
Heymann, 1972. Lunar soils show a totally unexpected concentration on grain size and by etching experiments to reside on grain surfaces by an inverse dependence of the which could only be produced by decay of U in the early packaging material [Leich et al., 1974]. Further (as was discussed in section B2), leaching [Tatsurnoto et al., 1973] and volatilization [Silber, 1974] experiments are best interpreted as indicating that Pb, rich in primordial $^4\text{Pb}$, is concentrated on the surfaces of the spheres. For more typical soil samples, studies of volatile Pb provide the only strong suggestion of surface deposits. The best case is for 14163 [Silber, 1972], where $\sim9\%$ of the lead evolved at temperatures less than 600ø.

Moreover, this Pb showed a high $^\text{206}\text{Pb}/^\text{204}\text{Pb}$ ratio ($\sim1.27$), which could only be produced by decay of U in the early history of the moon (before $4 \times 10^9$ yr ago). The low-temperature lead is required to be isolated from U for the past $\sim4 \times 10^9$ yr. This is most readily understood if the Pb is on the surface of the grains, consistent with its low release temperature. This is a somewhat startling result, given the opportunities for redistribution and mixing with more normal lunar Pb over $4 \times 10^9$ yr; however, it is consistent with the observation that the only volatile-rich lunar rocks from the highlands with ages of $3.85 \times 10^9$ yr or greater (section B2). It is very important to show whether this exotic Pb is actually lunar Pb over $4 \times 10^9$ yr; however, it is consistent with the low release temperature of the sample during meteorite impact or by postimplantation reactions at low temperatures, yielding released curves for the implanted $^4\text{Ar}$ qualitatively similar to those for the lunar $^4\text{Ar}$. Also, Bibring et al. [1974] obtained essentially the same release curves for CO for implantation energies corresponding to solar wind C and to atmospherically accelerated C, provided that the surfaces had first been made amorphous by irradiation with high doses of He ions. This suggests that the distribution of both the implanted and the reimplanted ions is controlled by the properties of the surface, particularly the amorphous layer, rather than by the implantation energy. This is reasonable when one considers the results on the H distribution discussed above.

Many studies of the chemistry of lunar surface layers have focused on reduction processes in order to understand the submicron metallic iron grains found in lunar soils by magnetic studies. Housley et al. [1974] have proposed that the fine-grained metal is formed by reduction of FeO by H in a soil sample during meteorite impact or by postimplantation reaction of solar wind hydrogen with oxidized iron, accompanied by diffusive loss of H$_2$O. Further evidence for metal produced by radiation is that metal particles from soils have W contents that are much larger than those for any type of meteoritic metal, although metal from an igneous fragment in 14321 also appeared to have a high W content. Tungsten behaves both as a lithophile and as a siderophile element; consequently, it is enriched in lunar rocks in comparison with other siderophile elements. Thus metal formed by reduction of lunar silicates would be expected to have high W, as is observed. Thus there is evidence that some larger metal fragments are formed by reduction, most likely by impact into H-bearing soils. However, it is also possible that fine-grained metallic Fe on surfaces could also be formed directly during the implantation process or by postimplantation reduction without the necessity of impact heating or melting. Using photoelectron spectroscopy, Yin et al. [1975] have observed...
metallic Fe formed during He bombardment of olivine at solar wind energies, suggesting that reduction during implantation does occur, although the chemistry of the reduction process is not obvious for the case of He bombardment.

Estimates of solar wind sputtering rates for lunar material range from 0.05 [McDonnell and Flavill, 1974] to 0.5 A/yr [Bibring et al., 1974]. The sputtering is primarily due to He ions because of the trade off between increasing sputtering yields and decreasing abundances for heavier ions. The above rates are much smaller than surface-averaged erosion rates of lunar rocks (~10 A/yr) measured from profiles of solar flare produced radioactive nuclei [Imamura et al., 1974] or from galactic cosmic ray track profiles [Behrmann et al., 1973]. Micrometeorites appear to be the chief erosional mechanism for lunar rocks, but for individual grain surfaces, sputtering appears to be the dominant erosional mechanism. Erosion by very small particles (nanometeorites) has not received much attention. In addition to erosion it is possible that sputtering can produce some of the interesting isotopic properties that appear to be characteristic of lunar surface layers. Figure 14 shows the isotopic composition of oxygen released by partial reaction of lunar soils as a function of the amount of oxygen liberated [Epstein and Taylor, 1973]. If the initial reaction steps preferentially remove material from the surface layers of the grains, the amount removed corresponds to ~50 A. The first fractions show very high depletions in 18O relative to 16O, which decrease rapidly toward typical bulk soil compositions as the surface layers are stripped away. Similar enrichments are observed for 34Si with respect to 28Si. Also, lunar soils appear to have systematically higher 40K/27Al [Barnes et al., 1973] and 35S/32S [Smith et al., 1973] ratios than lunar rock samples. However, the effects for K and S are 0.5-1% for a bulk sample and are surprisingly large in comparison with those observed for O and Si if they are due entirely to the surface layers. Further, a leaching experiment gave a 40K/44K ratio that was indistinguishable from the bulk soil, suggesting that the K isotopic variations are not a surface property; this is quite surprising in light of the O and Si results. Sputtering is only one possible explanation for the O and Si isotopic variations. Clayton et al. [1974] reject sputtering and prefer to ascribe the isotopic variations to vapor deposition and preferential loss of light elements in the lunar atmosphere.

Accretionary particles are very important lunar surface features, as is shown by dark surface coatings (patinas) of these on many rock surfaces [Blanford et al., 1974]. These are small, impact-produced, disk-shaped glass splashes of micron to submicron size that are formed in high abundance on many soil particles as well as on rock surfaces. They appear to be very fluid, and formation temperatures of 1300°-1700°C are estimated. These particles are to a certain extent a complication, because most of the interesting volatile elements are probably outgassed in the accretionary particle; thus they form a patch of fresh glassy surface and bury an older implanted surface in the process. Their presence leads to chemical and physical surface heterogeneities and complicates the interpretation of thermal release as well as many other 'traditional' experiments. It is just as well that these are cute little beasts, because we are going to have to learn to live with them. An interesting possibility put forward by Blanford et al. is that the highly outgassed accretionary particles could be the source of the observed O and Si surface isotopic variations.

In summary, the lunar surface layers at least are the site of very interesting processes and at best may represent a fundamentally different state of matter than has hitherto been available for study. It is also likely that similar radiation-damaged, implanted, amorphous material must characterize interstellar dust grains and possibly the grains that once populated our solar nebula at the beginning of the solar system. It
is also of interest to compare the properties of lunar surface layers with materials that are radiation damaged under laboratory conditions or conditions prevailing in the interiors of proposed fission and fusion reactors. I would argue that there are important problems as well as interesting science associated with the study of the outer lunar micron.

**Microcraters**

I find it difficult to explain to students why small, ≤1-cm-sized glass-lined impact pits on rocks were unexpected. Presumably, it was believed that impacts of micrometeorite particles would produce fracturing and spallation of rock surfaces without leaving visible craters. In any case, impact pits are now an accepted commonplace feature of lunar rocks and are discussed in detail in several reviews [Neukum et al., 1973; Hörz et al., 1975]. A crater of 0.05- to 1-mm pit diameter is characterized by a central pit with a depth to diameter ratio of ~0.7 [Brownlee et al., 1973], which is much larger than that for meter- or kilometer-sized craters. The central pit corresponds to material ejected from the immediate vicinity of the point of impact, possibly much of it being volatilized. This includes the projectile; no residues can be found in the bottom of the pits that can unambiguously be attributed to the micrometeorite producing the pit [Schneider et al., 1973]. The central pit is lined by a cup of glass with some evidence of flow at the margins. At greater distances from the point of impact there is insufficient energy imparted to the host material to cause melting but enough to cause mechanical displacement, leading to the formation of a distinctive ‘spall zone,’ characterized by an intersecting pattern of radial and concentric fractures. The ratio of spall to pit diameters is typically 4. For craters smaller than ~5 μm the spall zone is absent, and identification becomes more difficult [Neukum et al., 1972]. The observed microcraters are interpreted as being of ‘primary’ origin rather than as being produced by high-velocity ejecta from larger impacts (‘secondary’). The arguments for this are as follows:

1. In the required velocity range the flux of primary particles is many orders of magnitude higher than the flux of secondaries.

2. High-velocity secondary particles are preferentially emitted at low angles with respect to the lunar surface; thus microcraters from secondary particles should be concentrated on vertical faces of rocks rather than on horizontal surfaces, contrary to what is observed [Hartung et al., 1972].

These arguments may not be valid for micron and submicron craters. Figure 15 shows a composite crater frequency distribution compiled from data on several rocks and the corresponding mass of micrometeorites in a given pit diameter range. Most of the micrometeorite mass infall to the moon comes in microgram particles.

*Enhanced Iron Emission in Solar Flares*

The purpose of this section is to illustrate further the use of lunar samples and of terrestrial materials exposed in the lunar environment as detectors of particles from the interplanetary medium, the sun, and the galaxy. The preceding discussion of the use of microcrater densities to determine the long-term micrometeorite flux was one example. The present discussion concerns a result of great importance for cosmic ray physics and for the determination of solar system elemental abundances.

The most valuable artifact from Surveyor 3 returned by Apollo 12 was a filter from the TV camera, which because of 2.5-yr exposure to solar flares contained readily measurable densities of particle tracks (Figure 16) from the Fe group nuclei in the flares [Price et al., 1971]. As is illustrated in Figure 16, the track densities were greatly in excess of those calculated by using satellite solar flare He fluxes during the same period and by assuming that the Fe/He ratio in the flares was the same as that in the solar photosphere. Also, larger
of these results, e.g., the solar flare Fe enhancements, are now widely accepted and essentially established facts. Others, like the study of impact breccias and microcraters, are now mature fields of study with well-established bodies of data. In contrast, topics like the study of the regolith and the outer micron are immature research areas from which much is yet to be learned. Finally, results such as the ancient lunar magnetic field and the early intense bombardment have implications far beyond lunar science and when they are understood, will reshape our view of the solar system as a whole.

My conclusion is that lunar science, both past and future, and the carefully protected lunar rocks are the true legacies of the Apollo program. Hopefully, less biased historians of the future will concur.

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Because of the large number of references to the Lunar Science Conference proceedings we have adopted an abbreviation for the conference number and volume in order to save space. The first digit after the prefix LSC refers to the conference, and the second digit to the volume. For example, LSC3-1 refers to volume I of Proceedings of the Third Lunar Science Conference. The Apollo 12 and Third Conference proceedings were published by MIT Press. The other proceedings were published by Pergamon Press.


Epilog

In the preceding section, I have summarized a few of the exciting and unexpected results that were obtained from the Apollo program. The choice of topics is clearly personal. Some enhancements were observed at lower energies (shallower depths). This was subsequently confirmed by plastic track detectors flown on the Apollo 16 cosmic ray experiment [Bradley et al., 1973; Fleischer and Hart, 1973], which were exposed to the April 16, 1972, flare, and similarly by low-energy particles detected in plastics on the Apollo 17 cosmic ray package (R. T. Woods et al., unpublished manuscript, 1974), even though no visible flare activity occurred during Apollo 17. The Surveyor glass conclusions were further verified by detectors on satellites [Mogro-Campero and Simpson, 1972], which also showed that there is a systematic increase in the heavy element overabundance with increasing mass, demonstrating preferential acceleration of heavy nuclei in solar flares. Previously, it had been assumed that nuclei with the same nuclear charge to mass ratio would not be fractionated during acceleration and that the relative intensities of solar flare ions could be used to obtain refined values for solar abundances [Biswas and Fichtel, 1965]. The preferential acceleration of heavy nuclei is believed to show that during the early stages of acceleration the atomic charge is less than the nuclear charge; thus Fe and He are differentially accelerated. One possible mechanism is that the source region contains a much higher fraction of ionized Fe and other heavy elements than He and the Fe is preferentially extracted [Hirschberg, 1973]. Independent of the detailed explanation, it is now clear that much care is necessary in deducing solar abundances from relative intensities of solar flare ions.

Fig. 16. Measured densities of solar flare Fe tracks penetrating to a given depth in Surveyor 3 glass compared with predicted track density based on satellite Fe fluxes. This was the first evidence that heavy elements were preferentially accelerated in solar flares.
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