THE 1959-60 ERUPTION OF KILAUEA VOLCANO HAWAII

PETROLOGY OF THE KILAUEA IKI LAVA LAKE

GEOLOGICAL SURVEY PROFESSIONAL PAPER 537-B
Petrology of the
Kilauea Iki Lava Lake
Hawaii

By DONALD H. RICHTER and JAMES G. MOORE

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A petrologic and thermometric study of a
semienclosed body of slowly cooling basalt

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By Donald H. Richter and James G. Moore

ABSTRACT

The 1959 summit eruption of Kilauea Volcano filled the crater of Kilauea Iki to a depth of 365 feet with 50 million cubic yards of tholeiitic basalt. In the 3-year period since the formation of this lava lake, the U.S. Geological Survey core drilled four holes through the crust for petrologic and thermometric studies of a semiclosed body of slowly cooling basalt.

Between the end of the eruption in December 1959 and December 1962, the crust over the lake of molten lava had attained a thickness of more than 43 feet. Growth of the crust is attributable to the radiation and conduction of heat which is controlled primarily by the amount of rain falling on the lake's surface. Rate of crustal growth has decreased from about 4 feet per month during the first four months of the lake's history to an average rate of 0.34 foot per month between June and December 1962.

The rocks in the crust of the lava lake are porphyritic olivine basalts that have a wide range in phenocrystic olivine content. The olivine phenocrysts are set in a fine-grained, generally holocrystalline, intergranular groundmass of clinopyroxene and plagioclase with minor opaque minerals, apatite, hypersthene and glass. The base of the crust is characterized by a 5- to 6-foot-thick transient zone of crystallization that contains as much as 30 percent interstitial liquid at the crust-melt contact. Olivine-poor diabase segregations have been found at depths below 32 feet and constitute less than 1 percent of the total core examined.

The bulk density of the crust is almost entirely a function of the degree of vesiculation and increases from about 2.0 g per cc at the surface to 2.8 g per cc below a depth of 34 feet. Densities of nonvesicular glasses that contain a few olivine phenocrysts range from 2.6 to 2.9 g per cc. The grain size of the groundmass reaches a maximum within 15 feet of the surface, regardless of the thickness of the crust.

A total of 20 new chemical analyses together with 45 modal analyses of the rocks and melt in the upper part of the lake are presented. These data portray the chemical and mineralogical properties of the crust at five different times during the early history of the lake. The changes in composition in most of the rocks are due to two processes of fractional crystallization: gravity settling and filter pressing. Settling of olivine phenocrysts has influenced the composition of the entire crust, effecting changes in silica content between 45.61 and 49.61 percent.

The olivine-poor diabase segregations and the bottom-hole ooze found at depth in the crust are filter-pressed residual pore liquids and contain as much as 54.08 percent SiO₂. The low refractive index of the glass in the transient zone of crystallization indicates that differentiates with more than 60 percent SiO₂ are capable of being formed by filter pressing. The overall differentiation trend is toward pronounced enrichment in K₂O, Na₂O, and TiO₂ and moderate enrichment in total iron oxides with increasing SiO₂.

Chemical analyses of the melt indicate that a zone of transient alkali enrichment exists immediately below the crust. Enrichment of K₂O in this zone is upwards of 100 percent over that in the crust rocks and original lavas that filled the lake. The alkalis apparently moved from deep within the lake and concentrated in the melt below the crust. Upon crystallization of this zone, the excess alkalis were driven off and deposited as crystalline alkali sulfate sublimates in the cooler parts of the crust. It is postulated that, if analogous conditions exist in the magma reservoirs of the Hawaiian volcanoes, this process of alkali enrichment, coupled with removal of clinopyroxene by settling, could well explain the formation of alkali basalt from a tholeiite parent.

INTRODUCTION

The origin of the two principal basalt magma types—tholeiitic and alkalic—has been one of the most difficult problems in igneous petrology. In 1950, C. E. Tilley advanced our concepts of basalt petrogenesis by calling attention to the presence of tholeiitic magma in an oceanic environment and by demonstrating that a consistent chronological change in magma type, from tholeiitic to alkalic, occurred during the evolution of Hawaiian volcanoes.

Since 1950, the interest in the Hawaiian Islands has been attested to by the increased number of petrologic papers referring to Hawaiian basalts. As would be expected, however, the various investigators are not unanimous in their theories regarding the origin and relationship of the two principal basalt magma types. Unfortunately, deep-seated igneous processes cannot be
examined directly, but must be inferred from laboratory investigations of less complex systems and from field relationships. The inability to duplicate fully the physical-chemical conditions which control the natural systems leaves considerable room for speculation and argument.

The 1959 summit eruption of Kilauea Volcano has provided an unexcelled opportunity to study a slowly cooling body of tholeiitic basalt. At the end of the eruption in late December 1959, approximately 50 million cubic yards of molten lava filled the crater of Kilauea Iki to a depth of 365 feet. This exposed magma chamber, where processes of crystallization and magmatic differentiation are perhaps analogous to those within the Kilauea magma reservoir, presents an unparalleled opportunity for petrologic investigations. This ideal natural laboratory has already prompted a number of investigations. Shortly after the end of the 1959 summit eruption the staff of the U.S. Geological Survey's Hawaiian Volcano Observatory initiated a program of drilling and thermal measurement. Temperatures measured in the crust and melt of the lava lake during the early stages of this program have been the subject of two preliminary reports (Ault and others, 1961; Ault and others, 1962). The first hole to completely penetrate the crust of the lake was drilled in July 1960, by the Lawrence Radiation Laboratory (LRL) of the University of California, in an attempt to test drilling techniques in hot and molten rock masses (Rawson, 1960). Core and melt samples from the LRL hole have been studied and analysed by Macdonald and Katsura (1961). Ground magnetic surveys on the lava lake have been conducted by Decker (1962), and F. C. Frischknecht and L. A. Anderson (unpub. data) have measured the electrical conductivity of the melt by field electromagnetic techniques.

This paper (1) summarizes the physical, petrographic, and chemical data obtained from studies of core and melt, collected at known temperatures, from three successive U.S. Geological Survey holes drilled between April 1960 and June 1962, and (2) discusses the petrologic implications of these data. Although the crust of the lava lake was only a little more than 41 feet thick in June 1962, magmatic differentiation had already affected the composition of the rocks. In fact, the range in chemical composition in this relatively thin crustal layer is greater than any previously known for the exposed volcano of Kilauea.

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GEOLOGIC AND PETROLOGIC SETTING

Kilauea is one of five volcanoes that compose the Island of Hawaii, largest and southeasternmost island in the 1,500-mile-long chain of volcanic islands that form the mid-Pacific Hawaiian Archipelago (fig. 1). Kilauea, 4,090 feet above sea level, and its lofty companion Mauna Loa, 13,680 feet above sea level, are the youngest and most active of the Hawaiian volcanoes.

FIGURE 1.—Map of the island of Hawaii showing location of Kilauea Iki crater on Kilauea Volcano.
Both are typical shield volcanoes built by the outpourings of many thousands of relatively fluid flows of tholeiite basalt; neither volcano is known to have yet produced lavas of the alkali basalt suite, which are characteristically erupted intermittently and in minor volume during the dying stages of Hawaiian volcanoes.

The broad summit of Kilauea is indented by a large, relatively flat-floored caldera, approximately 3 miles long and 2 miles wide. In the southwest part of the caldera is Halemaumau, a precipitously walled 500-foot-deep crater with a diameter of about 3,200 feet. Two rift zones—a southwest and an east—extend down the flanks of the volcano, from the summit area to far below sea level. Summit eruptions, as the name implies, are restricted to the caldera region with most of them occurring in Halemaumau; flank eruptions, which generally follow a summit outbreak, occur along the rift zones.

Kilauea Iki, site of the 1959 summit eruption, is a deep and relatively large pit crater immediately east of the summit caldera. This crater is approximately 1 mile long and one-half mile wide, and its rim ranges from about 3,900 feet above sea level along the northeast and west sectors to about 3,600 feet above sea level along the low, narrow ridge that separates Kilauea Iki from the summit caldera (see fig. 2). The preruption floor of Kilauea Iki was a small flat plain, 37 acres in area, averaging 3,130 feet above sea level. This plain was the surface of a shallow lava pond formed during the brief eruption of 1888, the last eruptive activity in the crater prior to 1959.

The bulk of the tholeiitic lava flows of Kilauea is remarkably similar in composition, and any changes that do occur, both within the lavas of a single eruption and from eruption to eruption, are, in general, attributable to the addition or removal of olivine (Powers, 1955). The average silica content of Kilauea basalt, determined by 24 chemical analyses of prehistoric (pre-1790) and historic flows, is about 50 percent, the norm containing a few tenths of a percent olivine (Macdonald, 1949b, p. 74). The minor intrusive rocks and some of the massive flow rocks of Kilauea, on the other hand, show the effects of more advanced differentiation. Rocks from the Uwekahuna laccolith in the Kilauea caldera wall range from 45.7 to 52.04 percent in silica content, a range in composition as great as that found for all the analyzed Kilauea flows (Murata and Richter, 1961). Likewise, Kuno and others (1957) have found that segregation veins in a thick augite-olivine basalt flow exposed in the caldera wall contain 52.36 percent SiO₂, which, until the present study, represented the most siliceous tholeiitic differentiate found on Kilauea.

Recently, evidence has also been found suggesting that an increase in K₂O in the Kilauea magma has occurred during late prehistoric–early historic time (Moore and Richter, unpub. data). This change is in the direction of part of the shift from tholeiitic to alkalic basalt.

THE 1959 ERUPTION AND FORMATION OF THE KILAUEA IKI LAVA LAKE

The 1959 summit eruption of Kilauea Volcano began at 8:08 p.m. on November 14, 1959, when several discontinuous fissures halfway up the 600-foot south wall of Kilauea Iki began discharging liquid lava (Richter and Eaton, 1960). In 2 hours the line of eruptive fissures was approximately one-half mile long, with the individual lava fountains reaching heights of 100 feet. Through the night, however, activity gradually ceased, first along the outermost vents. By the next day only one vent, in the west end of the crater (vent, fig. 2), remained active. As the eruption continued both the rate of lava extruded from the single vent and the size of the lava fountain increased, with the fountain reaching heights of 1,200 feet by November 20. On November 21, after almost 7 days of continuous eruption, the level of the lava in the crater reached the level of the vent and the fountaining ceased.

In the next 4 weeks, 16 additional eruptive phases, ranging in duration from 32½ hours (4th phase) to 19½ hours (14th phase), occurred in Kilauea Iki. The 17th and last of the eruptive phases, stopped at approximately 8:00 a.m., December 20, 1959, marking the end of the summit eruption. All these later and shorter phases of the eruption were apparently fed by the same conduit-vent system which fed the week-long first phase, and, moreover, all erupting stopped soon after lake lava reached the level of the vent.

This phenomenon of lake lava drowning the active lava fountain and stopping the extrusion of lava was exceptionally well displayed during the eruption. As lake lava encroached upon the orifice of the vent, just prior to the end of an eruptive phase, the fountain activity became very erratic, suggesting that the relatively heavier, degassed lake lava was impeding the natural flow of new gas-charged lava from the vent. As the lake level continued to rise, it was apparent that great quantities of lake lava were actually pouring into the vent between bursts of fountaining, until finally, enough material filled the conduit to effectively prevent vesiculation and thereby pressurecapped the system. After the cessation of fountaining, the entire lake surface would immediately subside as more and more lake lava flowed back down the vent. This backflowing lava evidently returned to the magma reservoir from which it...
had risen, there to mix with new magma and again rise to the surface in a subsequent eruptive phase. Backflow often continued for as long as 2 days after the end of a phase, causing the lake level to subside as much as 50 feet. In the first hour of backflow the volume rate frequently exceeded 2 million cubic yards per hour, or more than twice the average rate of eruption.

The volume of lava erupted into Kilauea Iki during the 17 eruptive phases, and the volume withdrawn following each phase is shown graphically, as a function of time, in figure 3. A quantitative measure of the rate of extrusion and withdrawal is expressed by the slope of the lines in the graph. During the week-long first phase, the rate of extrusion increased from 40,000 cubic yards per hour for the first 2 days of eruption to a maximum of 500,000 cubic yards per hour on the sixth day and then decreased slightly to 300,000 cubic yards per hour on the last day. During the later phases, rates of extrusion (and withdrawal) were generally much higher. The maximum rate measured was 1,600,000 cubic yards per hour during the 3½-hour-long 16th phase.

A total of 40 million cubic yards of lava filled the crater at the end of the first phase; backflow was relatively minor and resulted in a loss of only 1,500,000 cubic yards. The very obvious change in the average net capacity of the lava lake (fig. 3) from 40 million to 50 million cubic yards between the third and fourth phases was due to a change in the elevation of the vent. The fourth phase activity started on the steep, inner
side of the new cinder-pumice cone approximately 40 feet higher but over the same area as the vent of the earlier phases. In all subsequent phases the vent remained at this higher but somewhat variable level allowing the lake to fill to a much greater depth before flowing back into the vent. Following five of the eruptive phases (Nos. 2, 7, 9, 12, and 15), more lava was withdrawn than had been erupted during each phase. Apparently some form of breakdown around the orifice of the vent, probably caused by the backflowing lava, allowed some of the lake lava from the earlier phases to escape. The lake attained its greatest depth (414 ft) and volume (58 million cu yd) at the end of the eighth phase on December 11, 1959. After the end of the last backflow on December 23, 1959, the lava lake contained slightly less than 50 million cubic yards and had a maximum depth of approximately 365 feet.

With the exception of the pyroclastic debris, most of which built the cinder-pumice cone leeward of the vent (fig. 2), all the lava extruded during the eruption was contained in Kilauea Iki. Because of the extreme fluidity of the lava and rapidity of filling, it is assumed that most of the lava was thoroughly mixed, and any possible correlation between specific layers of lake lava and lava from an individual eruptive phase is unlikely. Furthermore, since filling, convection currents have no doubt contributed to a certain amount of mixing within the still fluid part of the lake.

The filling of the crater was neither orderly nor systematic. In the first phase, most of the early lavas poured into the trough leading from the shallow west end to the deep east end of the crater and spread rapidly across the crater's flat floor (fig. 2). By the time lava in the bottom of the crater reached a depth of 25 feet, new lava flowing down the trough from the vent had begun to plunge under the crust of the lake, floating it upward and filling the young lake from below. Although this process continued throughout most of the first phase, the original crust on the lake did not persist more than a few days and, in fact, a

![Graph showing volume of lava erupted into Kilauea Iki during the 1959 summit eruption. Dashed line after the end of an eruptive phase represents backflow of lava down the vent. Each dot is a volume measurement. Note that the volume of total lava actually decreased during the fourth eruptive phase.](image-url)
new crust formed over most of the lake at least once or twice every 24 hours. As the crust floated upward, a zone of very thin and unstable crust was left between the thick, older crust and the sloping crater walls. From this peripheral weak zone, liquid lava continually welled up from below and spread out over the old surface, eventually cooling to form a new crust. In the later phases, most of the new lava contributed to the lake spread from the vent area as thick surface flows, covering and probably assimilating the old crust. In at least two phases, however, subcrustal injection played an important role in filling the lake.

The process of backflow following an eruptive phase further complicated the lake's history. Crustal fountaining, somewhat similar to that which occurred at the periphery of the lake during filling, continually reworked the lake surface during backflow. This reworking appeared to be a direct consequence of the liquid lake lava attempting to reach a common level in the concave bowl formed by the continual subsidence of the lake's crust. In this manner, an entirely new surface crust was generally formed every few hours on the lake. The end result on December 23, 1959, was an extremely level lake surface, with maximum relief of less than 5 feet over an area of approximately 135 acres (see section, fig. 2).

METHODS OF STUDY

In March 1960, less than 3 months after the end of the 1959 eruption, engineering surveys by the staff of the U.S. Geological Survey's Volcano Observatory were conducted in and around Kilauea Iki to set up controls for the various investigations planned on the lava lake. A 4,000-foot base line trending approximately N. 80° W. and a 2,000-foot crossline, both with permanent stations spaced every 100 feet, were laid out by transit and tape (fig. 2). Case-hardened 2-inch concrete nails driven into the crust through stamped aluminum tags were used to designate the permanent stations. Temperatures of more than 100° C within 2 inches of the surface were indicated by the fact that, after the nails were emplaced, water dropped on the nailheads would immediately boil.

After the control lines were installed, a level line was carried into the crater and the elevations of all permanent stations were determined. Releveling, repeated every 3 to 8 months since the original survey, has shown that the lake surface subsided an average of 0.7 foot per year during the 2-year period ending March 1962.

Access to the surface of the lava lake is by means of a 1-mile-long foot trail which switchbacks down the 400-foot-high east wall of the crater (fig. 2).

DRILLING AND SAMPLING

A total of four holes (fig. 2) has been drilled by the U.S. Geological Survey into the crust of the lava lake between April 1960 and December 1962. As mentioned previously, however, only data from the first three holes are fully presented in this paper. All holes were core drilled their entire length, but core recovery seldom averaged more than 20 percent, principally because of the extreme temperatures and the consequent unorthodox drilling methods used. Melt was collected below the crust in holes 1 and 3, and during the first penetration of hole 2; no attempt was made to sample the melt in the second penetration of hole 2.

On April 9, 1960, hole 1 was drilled to a depth of 7½ feet using an ordinary 1½-inch electric hand drill powered by a 300-watt portable generator. The bit employed was a hollow, tungsten-carbide masonry drill, 1½ inch in diameter and capable of holding 3 inches of core, welded to a section of 3½-inch solid steel rod. On April 25 the hole was deepened to 11.8 feet and on May 6 to 12.7 feet. Drilling was done dry to 7 feet; below that, water was sparingly poured into the hole during drilling in an attempt to cool the bottom. Despite the cooling, however, temperatures at depths of more than 12 feet often reached 900° C, and at 12.7 feet both the drill and bits failed to function properly and drilling was temporarily halted. Although the equipment used in this first drilling on the lava lake was primitive, it demonstrated that core drilling in extremely hot rock is feasible.

Hole 1 finally penetrated the crust on August 24, 1960, using a 1½-inch core drill powered by a 2½-hp gasoline engine with a maximum speed of 400 rpm. The drill bits, specially fabricated by Sprague and Henwood, Inc., consisted of tungsten-carbide cutters set in a high-temperature matrix with nickel shims. Core was collected in a conventional thin-wall, 2-foot double-tube core barrel. During drilling, water was manually pumped into the drill rods through a water swivel designed and built by Chester K. Wentworth of the U.S. Geological Survey.

Using the same drilling equipment, hole 2 was drilled through the crust (29.7 ft thick) on April 13, 1961, and again on October 4, 1961 (35 ft thick). Immediately after the second penetration in October, a stainless steel-ceramic probe containing a thermocouple was forced 4 feet into the melt below the crust (Ault and others, 1962).

Hole 3 was drilled through the crust (41.6 ft thick) between May 31, and June 11, 1962, again using the same equipment as that used for hole 2. An attempt to force a 10-foot probe into the melt proved unsuccessful, and the hole, still containing the probe and approximately 15 feet of drill rod, was abandoned.
With more elaborate and powerful drilling equipment (a conventional 9-hp portable drill and a 3½-hp water pump), hole 4 penetrated the melt at 43.6 feet on December 6, 1962. On January 10, 1965, after extreme cooling with water, hole 4 was deepened to 48 feet, or about 4 feet below the normal crust-melt boundary.

Drilling rates through the vesicular basalt of the crust could be maintained, under ideal conditions, as high as 1 foot per minute. In the denser rocks, below about 30 feet, drilling was generally slower; and within a few feet of the melt drilling was always very slow, with rates rarely exceeding 0.1 foot per minute. Moreover, extreme care had to be exercised when approaching the melt owing to liquid oozing from the wall of the hole and freezing onto the drill bit. On more than one occasion the bit stuck fast in the hole and the drill stalled. When this occurred, the rods and bit were left without benefit of coolant water until thermal equilibrium returned (generally 15 to 20 min) after which the string of rods could, with considerable difficulty, be withdrawn.

The drill bit, however, could not survive being submerged too long in the highly corrosive melt. When the melt was penetrated at 22.5 feet in hole 1, the bit became stuck and the rods snapped 15 feet below the collar of the hole. The rods were finally extracted after 1½ hours, during which time the entire bit plus 5 inches of the double-tube core barrel had been completely dissolved by the melt (fig. 4).

Drill rods were pulled after every 3 inches during the drilling of hole 1 when the masonry bit was used, and at least after every 2 feet when the conventional core barrel was used. Care was taken to reduce grinding up of the core to a minimum, but even with precaution, many drilling runs produced no core whatsoever. Melt was collected with either the drill bit or a specially designed 5-inch-long, hollow steel spoon. Samples of core from the three holes are numbered consecutively, with respect to depth, as follows: Hole 1, KI-100 to KI-115; hole 2, KI-150 to KI-186; hole 3, KI-200 to KI-236. (See figs. 6, 7, and 8.)

![Figure 4](image-url)  
**Figure 4.—Core barrel and bit after 1½ hours of immersion in melt at a depth of 22.5 feet. Compare with new core barrel and bit.**

**THERMOMETRY**

It is not the intent of this paper to discuss fully the temperature investigations that have been undertaken on the lava lake. However, the temperature profiles measured in the holes as soon as possible after drilling and hence, indicative of the sample temperature at time of collection, are important in understanding the crystallization history of the lava lake. Not all the many temperature profiles measured through the crust of the lava lake since April 1960 are used in this paper. These valuable source data are the basis for studies presently underway on the physical properties of the crustal rocks and for calculations of physical constants, such as thermal diffusivity and latent heat of crystallization.

Most of the temperatures obtained in the drill holes were measured with sheathed chromel-alumel thermocouples and a slide-wire potentiometer with a reading accuracy of 0.005 millivolt. The sheathed thermocouples consisted of matched chromel-alumel pairs, swaged with powdered MgO filler, in a thin-wall 310-stainless-steel case. Unsheathed thermocouples were used successfully for the first measurements in hole 1, but after melt penetration, a few seconds exposure to the extremely hot SO₂-rich bottom-hole vapors would disintegrate the alumel element. In hole 3 a compound 5-pair chromel-alumel thermocouple, with junctions spaced 3 feet apart, was used. Laboratory calibrations, using the melting-point technique and latterly a comparison with a National Bureau of Standards calibrated platinum-platinum-90, rhodium-10 thermocouple, indicate an accuracy of about one-half of 1 percent at 500°C and about 1 percent at 1000°C for the thermocouple temperatures.

Bottom-hole temperatures were also measured, whenever conditions permitted, with incandescent filament-type optical pyrometers. In the shallower holes (hole 1 and first penetration of hole 2), optical-pyrometer temperatures were within 5°C of those obtained by chromel-alumel thermocouples. In the deeper holes, however, a greater divergence was observed, with the optical-pyrometer values consistently higher by as much as 10°C. Surface and near-surface (less than 1 ft) drill-hole temperatures were measured with mercury thermometers.

**PHYSICAL PROPERTIES AND PETROCHEMISTRY**

Rock-density determinations were made on most uniform pieces of core greater than one-half inch long. The ends of the selected core samples were ground flat to form a near-perfect cylinder, and the dimensions (length and diameter) were measured with vernier...
calipers. The weight of the individual cylinders was determined on a laboratory balance.

A total of 21 samples, including rock, melt, and one clinopyroxene separate, were chemically analyzed in the Denver Rock Analysis Laboratory of the U.S. Geological Survey. The clinopyroxene from a diabase segregation in hole 3 was separated by means of heavy liquids and an isodynamic separator. Norms of the analyses were calculated by computer. Modes were determined for 45 samples, and with only a few exceptions, more than 50 points were counted per thin section. The composition of the olivines was determined by the X-ray method of Yoder and Sahama (1957), and the composition of the feldspars, where practical, by the method of Tsuboi.

GROWTH OF LAVA LAKE CRUST

In the 3-period between the end of eruption in December 1959 and the last drilling in December 1962, the crust of the Kilauea Iki lava lake attained a thickness of 43.6 feet. Although the growth of the crust has been studied only in a very restricted area over the deepest part of the lake (fig. 2), the level surface indicates a comparable growth rate—hence uniform thickness—over a much larger area. At the periphery of the lake and in the relatively shallow west end, however, the crust is presumably thicker.

The criterion used to identify the base of the crust when drilling was the depth at which drill rods noticeably sank into the melt. Even though this depth is a complex relation of several physical and mechanical factors (see Ault and others, 1962, p. 2811), temperatures measured in all the drill holes at this depth were consistently between 1057° and 1065° C. The base of the crust was also identified independently during the emplacement of the melt probe in hole 2 as the 1065° C isotherm (Ault and others, 1962), corroborating the temperatures measured at the base of crust determined by drilling. As would be expected, there is no sharp demarcation between solid and liquid; instead, the contact is a zone only a few inches thick of rapid change in the proportion of crystals and in viscosity. Thin sections of crust and melt across this narrow zone reveal a glass (liquid) content of 24 to 26 percent in the lowermost crust and 30 to 73 percent in the upper part of the melt. The relatively rapid change in degree of crystallinity over an exceedingly limited temperature interval (<5° C) is due to the onset of plagioclase crystallization at or slightly above the 1065° C isotherm. It should also be noted that the thermal profiles (table 1 and figs. 6, 7, 8) show that the base of the crust is marked by only a subtle change in the slope of the thermal gradient.

The growth of the crust is shown graphically in figure 5 as a function of time. The rate of growth decreased from an average of 2.8 feet per month, during the first 8.1 months, to an average of 0.34 feet per month during the 5.6 months ending December 6, 1962. In the 22-month period between August 1960 and June 1962, however, the rate of growth decreased only slightly from 0.95 to 0.80 feet per month. Five selected temperature profiles, based on measurements made when equilibrium returned after drilling and therefore representative of the core-sample temperature at time of collection, are listed in table 1. These same data are plotted graphically in figures 6, 7, and 8. A contoured temperature diagram of depth versus time for the crust of the lava lake is shown in figure 6. The descent of all isotherms below 600° C has been virtually linear, the rates ranging from 0.12 feet per month for 100° C to 0.6 feet per month for 600° C during the period between April 1960 and June 1962. Isotherms above 600° C were relatively steep during the early period of crust formation and since about January 1962 have shown a marked tendency to flatten.

The transfer and loss of heat from the upper part of the lava lake, which directly affects the growth of the crust, are controlled principally by (1) the interrelated effects of radiation and conduction and (2) the vaporization of water (rain). Although all these processes will be effective until thermal equilibrium between the lake and its environment is attained, each plays a dominant role at various periods in the lake's development. During eruption and in the first few hours of stable crust formation, loss of heat was almost entirely due to radiation. However, as the crust...
Table 1.—Kilauea Iki lava lake temperature profiles, holes 1 to 3

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<tr>
<th>Depth (feet)</th>
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1. Mercury thermometer.  
2. Optical pyrometer. All other temperatures by chromel-alumel thermocouple.  
3. Drilled 0 to 12.8 ft between Apr. 9 and May 6, 1960; drilled 12.8 to 22.7 ft on Aug. 24, 1960.

(a poor heat conductor) became thicker, radiation became less and less efficient and heat loss became dependent on the amount of heat transferred to the surface by conduction. A third cooling process, vaporization of water, is a function of the amount of rain falling on the lake and being changed to vapor upon contact with hot rocks in the crust. Because an average of approximately 100 inches of rain per year falls on the Kilauea Iki area, the vaporization of water is now by far the most important single process contributing to the cooling of the lava lake.

The amount of heat (heat of vaporization) required to change 1 gram of water at 100 °C from a liquid to a vapor is 540 calories. Assuming that the basalt has a specific heat of 0.20 and a density of 2.7 g per cc, 1 cc of basalt will have a thermal capacity (mass×specific heat) of 0.54 cal per deg C, or 0.54 cal will be released for every degree cooled. Hence, the amount of heat necessary to vaporize 1 cc (1 g) of water is equivalent to the heat released in cooling 1 cc of basalt from 1100 °C to 100 °C.

However, these calculations can only be used as a rough qualitative measure of crustal growth because they fail to consider processes that add heat to the crust. For example, if all the rain falling on the lake was utilized in cooling and no heat was added, approximately 8 feet per year of lake lava would be transformed into crust having a temperature of 100 °C. This is a rate of 0.67 foot per month, virtually double the observed rate (0.36 ft per month) between June

![Figure 6.—Isotherms in crust of Kilauea Iki lava lake plotted as a function of time. Diagram compiled from thermal data from drill holes 1, 3, and 4.](image-url)
and December 1962, and yet this rate does not include the contribution due to heat loss by conduction and radiation. The physical-chemical processes that add heat heat to the crust in significant quantities are (1) heat of crystallization, (2) solar radiation, and (3) convection. Heat of crystallization is by far the most important of these processes, especially in the lower 5 feet of crust where upwards of 70 percent of the total volume is still liquid. The heat contributed by solar radiation to the surface of the lake and by convective overturn in the melt are of secondary importance but need to be considered in any attempt to understand the heat budget of the lake.

Furthermore, the questionable validity of the basic assumptions (density and specific heat of the basalt) could also affect the calculation of the rate of heat loss due to cooling by water. If either the density or specific heat are more than assumed, then by definition the thermal capacity of the basalt would be correspondingly larger and hence more water would be required to cool an equivalent amount of basalt. On the other hand, recycling of the water in the crust could substantially add to the effectiveness of cooling by the vaporization of rainfall. In fact, it is entirely possible that much of the 100 inches of rain per year is used over and over again, especially as the crust thickens leaving a relatively cool top to condense the vapors rising from the hotter zones at depth.

Though the above considerations are somewhat speculative, the rate of heat flow through the crust of the lake may be compared with the rate of heat loss due to the vaporization of rainfall. On the basis of a uniform heat conductivity of $5.1 \times 10^{-3}$ cal per cm per deg C for the crustal basalt, the rate of heat flow in calories per second can be expressed by the equation:

$$\text{Rate of heat flow} = k \cdot T_1 - T_2 \cdot x \cdot A,$$

where $k$ is the thermal conductivity of basalt, $T_1 - T_2$ the difference in temperature between the top and base of crust ($1000^\circ$ C), $t$ the thickness in centimeters, and $A$ the area (1 cm$^2$). As all parameters are fixed, with the exception of $t$, it is apparent that the rate of heat flow varies inversely with thickness. Solving the equation for thicknesses of 10, 20, 30, and 40 feet (305, 610, 914, and 1,219 cm), the corresponding heat flow rates are 16.4, 8.1, 5.5, and $4.1 \times 10^{-3}$ cal per sec. Hence, for a crust thickness of about 40 feet, the heat loss due to vaporization of rainfall begins to exceed the heat loss by conduction.

THE CRUST

Megascopically, the bulk of the rocks in the 42-foot thick crust of the Kilauea Iki lava lake is porphyritic olivine basalts that have a wide range of content of phenocrystic olivine. The base of the crust is characterized by a 5- to 6-foot-thick zone of fluid-rich basalt, referred to as the transient zone of crystallization, which moves downward at a rate equal to the crustal growth rate. Olivine-poor diabase segregations found only at depths below 32 feet constitute less than 1 percent of the total crust thickness examined.

The basalts exposed on the surface of the lake and in cracks penetrating a few feet below the surface are highly vesicular, the voids constituting up to 40 percent by volume of the rock. The groundmass matrix is generally dense, breaks with a rough hackly fracture, and, depending on the degree of crystallinity, ranges in color from gray to nearly black.

VESICULARITY, DENSITY, AND GRAIN SIZE

The size, shape, and relative abundance of the vesicles in the crust of the lava lake are shown diagrammatically in the drill-hole sections in figures 7, 8, and 9. The extreme top of the crust, too thin to portray in the sections, is characterized by a chilled glassy skin not more than 1 inch thick, containing abundant vesicles and covering, in many places, large flat gas blisters (macrovesicles) as much as 3 feet in diameter. Immediately below the glassy skin the vesicles are generally more uniform in habit and, as would be expected, decrease in size and abundance with depth. With the exception of a number of anomalous dense layers in the upper part of the crust, the spherical vesicles decrease from 8 to 10 mm in diameter near the surface to 1 mm or less in diameter below about 34 feet. In addition to the spherical vesicles, relatively large flattened voids and irregular tubes, probably formed under conditions of subtle shearing within the crust, are locally common at depths below 26 feet. Vesicles in the diabase segregations are even more irregular and form intricate branching tubes as much as three-fourths inch long.

The thin dense layers restricted to the upper 14 feet of crust are apparently partially remelted remnants of old crust that were engulfed by liquid lava during the last period of crustal foundering on the lake's surface. Although probably limited in areal extent and erratically distributed, these dense zones have locally affected the composition of the overlying strata. In every case where a dense zone has been recognized it formed an impermeable barrier to sinking olivine crystals and was overlain by layers of crystal mush with as much as 37 percent modal olivine.

The bulk density of the crust is almost entirely a function of the degree of vesicularity; the effect of changes in the olivine content on the density is possibly only apparent, at depth, in hole 3 (figs. 8 and 9). Bulk densities range from a low of 1.8 to 2.1 g per cc near the
PETROLOGY OF THE KILAUEA IKI LAVA LAKE

MODAL COMPOSITION, IN PERCENTAGE BY VOLUME

![Diagram showing modal composition in percentage by volume.]

Figure 7.—Composite section through drill hole 1 with pertinent mineralogical and physical data. Relative size, shape, and abundance of the vesicles in drill core are portrayed diagrammatically by pattern. Dashed lines denote obvious textural and chemical discontinuities.

surface to a maximum of 2.7 to 2.8 g per cc below a depth of 34 feet. The increase from 1.8 to 2.5 g per cc occurs fairly rapidly and consistently in about the first 6 feet of crust; below that the increase from 2.5 to 2.8 g per cc is erratic and gradual. The anomalously low-density zone between 26 and 32 feet in hole 3 may be due to a corresponding low olivine content in this same general interval (fig. 9).

Bulk densities were determined for a number of nonvesicular glasses that were collected while still liquid. Melt samples, containing a few olivine phenocrysts, had densities of 2.66 g per cc (sample KI-186, 35.0 to 35.2 ft, hole 2) and 2.78 g per cc (sample KI-235, 42.05 to 42.3 ft, hole 3). The silica-rich liquid that oozed into the bottom of hole 1 (sample KI-113, 22.3 to 22.5 ft) from the base of the crust had a density of 2.65 g per cc. Moreover, the specific gravity of vesicle-free fragments of tachylite from the surface of the lake ranges between 2.6 and 2.9. H. A. Powers (written commun.) has observed that vesicle-free glass shards from the Kilauea Iki lavas had specific gravities very close to 2.9 or perhaps even more.

The groundmass grain size (figs. 7, 8, and 9) does not show a consistent increase with depth as might be expected. There is an increase in size directly below the surface and directly above the base of the crust, but the maximum grain size appears to be attained within 15 feet of the surface, regardless of the thickness of crust.

PETROCHEMISTRY

A total of 20 new chemical analyses together with 45 modal analyses of the rocks and melt in the crust of the lake are presented in tables 2, 3, 4, and 5. These data portray the chemical and mineralogical properties of the crust at five different times (Apr. 9 to May 6, 1960; Aug. 24, 1960; Apr. 13, 1961; Oct. 4, 1961; and May 31 to June 11, 1962) during the early history of the lake. Although melt penetration occurred on the four latter
dates, only the samples collected on April 13, 1961, and May 31 to June 11, 1962, represent isochronal surface to melt profiles. The dates of sampling and the locations of the samples are shown on the diagrammatic drill-hole sections in figures 7, 8, and 9.

The chemical analyses were done by standard gravimetric methods in the Denver laboratories of the U.S. Geological Survey by the analysts credited in tables 2, 3, 4, and 5. All these analyses appear consistent; that is, they plot on smooth linear control lines in variation diagrams when compared with one another, with analyses of Kilauea lavas earlier than those of the 1959 eruption, and with Kilauea Iki analyses made by other laboratories (Tilley, 1960). Only the 10 Kilauea Iki analyses of Katsura (in Macdonald and Katsura, 1961, table 1, p. 362) show a marked divergence from the analyses presented here; Katsura’s analyses when plotted on a silica-variation diagram, similar to that used in this paper (fig. 10), are erratic and define, only within broad limits, the linear control lines tightly controlled by analyses from other laboratories. Also, Katsura’s analyses show a bias (again comparing with analyses from other laboratories) toward low silica by as much as 2 percent. The low silica values are partly compensated for by anomalously high alumina values, and it is the writers’ opinion that the apparent discrepancies lie largely in the values reported for these two constituents.
OLIVINE BASALT

In thin section the predominant tholeiitic olivine basalt samples consist of large olivine phenocrysts set in a groundmass of fine-grained clinopyroxene, plagioclase, and opaque minerals with varying amounts of glass (see fig. 11). Minute needlelike crystals of apatite (?) are common in the glass of the more vitric rocks, and minor hypersthene was observed in a few thin sections. The groundmass texture grades from completely glassy at the surface, through interstitial in a zone a few feet below the surface, to intergranular below about 5 feet. Beginning in the transient zone of crystallization at the base of the crust and going upwards, this textural sequence is reversed with the glass representing a true interstitial liquid phase at the temperature and time of collection.
The extreme variation in chemical composition of the olivine basalts is controlled principally by the physical addition or removal of olivine phenocrysts. The content of modal olivine ranges from 2 percent (sample KI-184) to 37 percent (sample KI-109), effecting changes of more than 4 percent in silica and more than 15 percent in magnesia. This pronounced olivine control (Powers, 1955) on the composition of the basalts is best demonstrated by the silica-variation diagram in figure 10. Each of the major constituent oxides, when plotted against SiO₂, defines a unique straight line between about 50 percent SiO₂ and the plot for the composition of hypothetical olivine (Fa₃₅) at 40 percent SiO₂. Rocks containing more than about 50 percent SiO₂ and less than 7 to 8 percent MgO are, in general, olivine poor and hence are unaffected by olivine control.

Although the modes of the groundmass minerals in the olivine basalts vary widely depending on the concentration of phenocryst olivine, the relative proportions of these minerals in the glass-poor basalts are fairly constant. The most striking constancy of volume ratios is between clinopyroxene (37 to 65 mode percent) and plagioclase (19 to 33 mode percent), which, with only a few exceptions, is remarkably close.
### Table 3.—Chemical analyses, norms and modes of rocks from hole 2, Kiluaea Iki lava lake

[Results in percent. Analyses by D. F. Powers, except samples KI-184 and KI-186 by C. T. Parker]

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#### Norms

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<th>ab.</th>
<th>an.</th>
<th>di.</th>
<th>wo.</th>
<th>en.</th>
<th>bi.</th>
<th>fa.</th>
<th>ho.</th>
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<td>18.80</td>
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<td>19.67</td>
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| Total | 99.87 | 99.99 | 100.00 | 99.99 | 100.03 | 99.96 | 99.95 | 100.01 | 100.09 | 100.05 | 99.99 |

#### Modes

| Phenocrysts: Olivine... | 15 | 5 | 29 | 19 | 28 | 7 | 23 | 32 | 5 | 8 | 14 | 7 | 11 | 6 | 8 | 20 | 2 | 6 | 7 |

| Groundmasses: | Pyroxene... | 47 | 53 | 41 | 49 | 44 | 59 | 46 | 60 | 65 | 56 | 55 | 55 | 50 | 50 | 45 | 42 | 27 | 50 | 43 | 43 |

Phlogopite... | 21 | 20 | 24 | 20 | 22 | 25 | 24 | 20 | 20 | 24 | 24 | 23 | 24 | 24 | 24 | 21 | 21 | 17 | 18 | 24 | 21 |

Opacites... | 9 | 8 | 9 | 7 | 9 | 8 | 9 | 11 | 12 | 7 | 7 | 7 | 5 | 4 | 3 | 5 | 5 | 3 | 2 | 4 |

Glases... | 12 | 9 | 1 | 4 | 1 | 5 | 2 | 0 | 2 | 1 | 1 | 0 | 8 | 10 | 17 | 24 | 30 | 41 | 13 | 22 | 26 |

Apatis... | Tr. | Tr. | Tr. | Tr. | Tr. | Tr. | Tr. | Tr. | Tr. | Tr. | Tr. | Tr. | Tr. | Tr. | Tr. | Tr. | Tr. | Tr. | Tr. | Tr. | Tr. |

---
1 Probably includes some opaques.
THE 1959–60 ERUPTION OF KILAUEA VOLCANO, HAWAII

Table 4.—Modes of rocks from hole 3, Kilauea Iki lava lake

<table>
<thead>
<tr>
<th></th>
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<tr>
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<td>8–9</td>
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<td>14–15</td>
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<td>21–22</td>
<td>24–25</td>
<td>27–28</td>
<td>29–31</td>
<td>33–34</td>
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<td>Tr.</td>
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<td>6</td>
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<tr>
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<td>30</td>
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<td>18</td>
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<tr>
<td>Plagioclase</td>
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<td>22</td>
<td>22</td>
<td>24</td>
<td>23</td>
<td>26</td>
<td>28</td>
<td>29</td>
<td>32</td>
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<td>31</td>
<td>25</td>
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<td>Opaques</td>
<td>10</td>
<td>11</td>
<td>8</td>
<td>5</td>
<td>12</td>
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<td>7</td>
<td>8</td>
<td>10</td>
<td>7</td>
<td>8</td>
<td>8</td>
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<td>1</td>
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<td>0</td>
<td>2</td>
<td>1</td>
<td>Tr.</td>
<td>0</td>
<td>15</td>
<td>Tr.</td>
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<td>Apatite</td>
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</tbody>
</table>

Figure 10.—Silica-variation diagram of all analyzed crust rocks and melt samples from Kilauea Iki lava lake. Calculated average composition of Kilauea Iki lava (K. J. Murata, unpub. data) is shown by + at 48.21 percent SiO₂. Lines drawn through the oxide plots below 49.61 percent SiO₂ represent olivine (Fa₃₃) control lines. Note change in scale for total iron.
to 2:1. In the olivine basalts with more than about 20 to 30 mode percent interstitial glass, on the other hand, there is a marked increase in the volume ratio of clinopyroxene to plagioclase. Melts collected at the base of the crust contain, after quenching, between 30 and 74 mode percent glass, and the volume ratio of clinopyroxene to plagioclase is consistently greater than 2.5:1 and is as high as 4.5:1 (sample KI-236). These relations suggest that clinopyroxene begins to crystallize at somewhat higher temperatures than plagioclase in the melt. The olivine phenocrysts in the crust range in size from 0.08 to over 4 mm in diameter. In the upper 2 to 3 feet of crust, olivine is euhedral to slightly rounded; below 3 feet, however, and throughout the remainder of the crust, it is highly rounded and embayed due to resorption by the melt. Evidence of reaction between the olivine phenocrysts and melt is spotty and inconsistent. Abnormal concentrations of clinopyroxene around the margins of the olivine were observed below a depth of 27 feet in hole 2 and below a depth of 5 feet in hole 3. Yet in hole 1 no apparent coronas were evident. Small crystals of hypersthene (?) were present on resorbed olivine crystals at a depth of 2 to 2.5 feet in hole 3 (sample KI-203) but were not observed to bear this relation elsewhere in the crust. The composition of five olivine separates from depths of 2 to 2.5, 8 to 9, 17 to 18, 29 to 31, and 37.7 to 39 feet in hole 3 ranged from Fa14 to Fa17. No consistent relation between composition and depth is apparent, and the olivine crystals show no evidence of zoning.

Clinopyroxene, the most abundant groundmass mineral in the olivine basalts, occurs as small, stubby hypidiomorphic prisms and irregular grains averaging 0.05 mm in diameter. Sparse microphenocrysts as much as 0.6 mm in diameter and with ragged crystal outlines were observed scattered through the groundmass in a few sections at a depth of more than 14 feet in holes 2 and 3. A tendency toward a subophitic intergrowth is discernible in some of the sections, but in general the clinopyroxene crystals are smaller than the plagioclase laths. This is a rather curious feature inasmuch as the clinopyroxene begins to crystallize at slightly higher temperatures and has a higher ratio to plagioclase in the glassy rocks than in the holocrystalline rocks. Glomerocrystic clots of clinopyroxene and plagioclase are common in the vitric basalts at the surface of the crust but are absent in the glass-rich basal zone. In this basal zone and the underlying melt the clinopyroxene is present as minute perfect euhedra dispersed through the glass. Zoning is not apparent in either the microphenocrysts or groundmass clinopyroxenes, but the latter do exhibit simple twinning. The groundmass clinopyroxenes are augitic with an approximate composition of $\text{Wo}_{0.2}\text{En}_{0.8}\text{Fs}_{0.10}$ (large 2V, $\gamma=1.710$); no optical properties indicative of pigeonite were observed. Clinopyroxene with $\text{Fs}\text{En}$ approximately $17:83$ is apparently in equilibrium with olivine $\text{Fa}_{14}$ to $\text{Fa}_{17}$.

The plagioclase occurs as hypidiomorphic laths, plates, and needles as much as 1 mm in length. In the clinopyroxene-plagioclase glomerocrysts typical of the vitric basalts at the top of the crust, the plagioclase commonly occurs as radiating star-burst clusters of laths, each lath mantled by a concentration of small clinopyroxene crystals. Slight zoning was observed in some of the larger crystals. The composition of the plagioclase at a depth of 34.3 to 35 feet in hole 2 (sample KI-186) is $\text{An}_{44}$.

Hypersthene is rare in the basalts and was never observed to exceed more than 1 percent by volume of the rock. Small crystals, probably of hypersthene, were observed on resorbed olivine phenocrysts near the surface in hole 3 (sample KI-203), but elsewhere hypersthene forms relatively large (as much as 0.1 mm) irregular to blocky prisms scattered through the groundmass. In the modal analyses (tables 2, 3, and 4) the minor amount of hypersthene is included in modal pyroxene.

Opaque minerals occur in a variety of forms in the groundmass and constitute between 3 and 12 percent of the total volume of the rock. Within 10 feet of the surface the opaque minerals generally form irregular masses with abnormally high concentrations around the periphery of the vesicles. At depths greater than about 10 feet they tend to be more regular in outline, forming rods, equant crystals, and chainlike groups of equant crystals as much as 1 mm in length. Although no large crystals of opaque minerals were observed in the melt, the general opacity of the melt glass is probably due in part to included opaque microcrystals. All of the opaques appear to be strongly ferromagnetic and hence rich in magnetite. The high titania content of the basalts, however, strongly indicates the presence of ulvospinel $(\text{FeTiO}_3)$ molecule in the magnetite and probably primary ilmenite as well.

Minute needlelike crystals of apatite (?) occur in trace amounts throughout much of the glass and are especially common in the glass-rich basalt in the transient zone of crystallization. The needles occur singly and in tufted groups.

Cristobalite was first identified in cores from hole 4 drilled December 3 to 6, 1962. It was not observed
in any core from the earlier drill holes. The cristobalite occurs as well-formed white pseudoisometric crystals as much as one-half mm in size lining vesicles. It occurs in much of the core in the interval 11.9 to 15.9 feet and also near the bottom of the interval 15.9 to 19.9 feet. Some of the rock probably contains up to 1 percent cristobalite, and all the rock containing the mineral is otherwise rather normal-appearing olivine basalt with 10 to 20 percent olivine. The temperature range of the cores containing the cristobalite was 400° to 615° C at the time of collection, and hence the cristobalite crystallized at a higher temperature range. Because it occurs only in vesicles in olivine-rich rocks, the cristobalite must have crystallized from hot gases streaming through the crust from the melt below. Such gas transfer indicates depletion of SiO₂ in the melt and enrichment of SiO₂ in the crust.

The glass content decreases from about 80 mode percent in the rapidly chilled upper inch of crust to 0 to 3 mode percent below a depth of 5 feet (figs. 7, 8, and 9). About 6 feet above the base of the crust, at the top of the transient zone of crystallization, the glass content again increases and reaches approximately 30 mode percent (sample KI-168) at the crust-melt contact. Most of the glass in the upper part of the crust is dark brown to nearly opaque, typical basaltic glass (n=1.61 to 1.63) probably corresponding to compositions of 49.02 to 49.85 percent SiO₂ and 7.44 to 10.20 percent MgO (K. J. Murata, unpub. analyses of glass from 1959 eruption). The low-silica and high-magnesia content of some of these glasses indicates that they would have crystallized significant amounts of magnesian olivine if they had not been quenched. The glass in the lower part of the crust and melt is discussed in more detail in the following sections.

**TRANSIENT ZONE OF CRYSTALLIZATION**

Although the rocks in the zone of crystallization are virtually all olivine basalts, similar mineralogically and chemically to the overlying holocrystalline basalts, the unique character of the zone and significance of the included magmatic fluids warrants a separate description.

The base of the transient zone of crystallization is the crust-melt contact. Here, as previously described, the transition between crust (<30 percent liquid) and melt (>30 percent liquid) is relatively abrupt and occurs within a thin zone not more than a few inches thick at about the 1065° C isotherm. The less well defined top of the zone, where pore liquid ceases to exist, lies 5 to 6 feet above the base of the crust at approximately the 970° C isotherm. As the crust of the lake grows, the zone of crystallization will not only advance downward but will also increase in thickness owing to a decrease in the lake's thermal gradient.

The transient nature of the zone of crystallization is well illustrated in the two photomicrographs in figure 11. Figure 11A depicts the state of crystallization within the crust-melt contact (sample KI-168) at a depth of 29.7 to 29.9 feet and at a temperature of 1065° C (fig. 8) in hole 2 on April 13, 1961. The abundant glass, which constitutes 30 percent of the rock (table 3), pervades the entire groundmass and surrounds the crystals and crystal groups of clinopyroxene, plagioclase, and opaques. Figure 11B shows a thin section of core (sample KI-225) from the same depth (29 to 31 ft), but collected about 14 months later (June 1, 1962) in hole 3 at a temperature of approximately 825° C (fig. 9). Crystallization is virtually complete, and only a trace of glass (table 4) remains in the groundmass. Moreover, the amount of opaque minerals has about doubled (from 3 percent in sample KI-168 to 7 percent in sample KI-225) during the course of crystallization.

Unlike the glass in the upper part of the crust, which was quenched, the glass in the transient zone of crystallization represents a sample of residual pore liquid in equilibrium with the groundmass minerals at the temperature of sampling. Hence, the composition of these glasses is strongly dependent on temperature and exhibits marked changes through the relatively narrow zone of crystallization. The relationship between volume, temperature, and index of refraction of the glass in the base of the crust on October 1961 and June 1962 is shown graphically in figure 12. Upward from the base of the crust the index of refraction of the glass decreases, concomitant with a decrease in amount and temperature, from 1.61 (melt, sample KI-236) to 1.508 (sample, KI-231), 4½ feet above the base. The color of the glass also changes through this interval from dark brown at the base to very pale brown. A very crude estimation of the composition of these glasses can be made by reference to the work of Stewart (1962) on the indices of refraction of fused Hawaiian rocks. Analyses showed that the melt glass with n=1.61 corresponds to a composition of about 49 percent SiO₂, in good agreement with the assumed compositions for glass of similar index at the surface of the crust. Using Stewart's least-squares curve showing the relation between index of refraction and SiO₂ content for all rocks studied (both alkalic and tholeiitic), we found that the lowest index of refraction (1.508) observed corresponds on the same curve to 59 percent SiO₂. However, because of a possible pronounced divergence between the
alkalic and tholeiitic index of refraction curves with increasing silica content, the low index glass may contain somewhat more than 59 percent SiO₂.

Two pore fluids from the base of the crust have been analyzed (table 5). One is the diabase segregation (sample KI-228) described in the following section, and the other is a glass (sample KI-113) collected from the bottom of hole 1 (22.3 to 22.5 ft). The glass (n=1.569, Stewart, 1962) represents a low-viscosity liquid that oozed from the base of the crust into hole 1 after the August 24, 1960, drilling and chanced to adhere to the end of a thermocouple used during a temperature-profile measurement. The glass contains 54.08 percent SiO₂ and 5.02 percent total alkalis—the most silica- and alkali-rich tholeiitic material ever analyzed from Kilauea. Neither the diabase segregation nor the glass ooze, however, are appreciably enriched in iron. The relative silica content of the two differentiates indicates that the ooze was derived from a higher level in the transient zone of crystallization and at lower temperatures than the segregation.
## TABLE 5.—Chemical analyses, norms, and modes of tholeiitic differentiates Kilauea Iki lava lake

(Results in percent. D. F. Powers, analyst)

<table>
<thead>
<tr>
<th>Sample and depth (feet)</th>
<th>KI-113 (ovine, hole 1)</th>
<th>KI-228 (diabase segregation, hole 3)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>22.3-22.5</td>
<td>33-34</td>
</tr>
</tbody>
</table>

### Chemical analyses

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<tr>
<th>Element</th>
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<th>Sample KI-228</th>
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</thead>
<tbody>
<tr>
<td>SiO₂</td>
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<td>50.32</td>
</tr>
<tr>
<td>Al₂O₃</td>
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<td>13.35</td>
</tr>
<tr>
<td>P₂O₅</td>
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<td>10.39</td>
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<td>MgO</td>
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</tr>
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<td>0.00</td>
</tr>
<tr>
<td>TiO₂</td>
<td>3.37</td>
<td>4.40</td>
</tr>
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<td>P₂O₅</td>
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<td>F⁻</td>
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</table>

Subtotal: (1) 100.11
Less O: 100.03
Total: 100.08

### Norms

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<th>Norm</th>
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<th>Sample KI-228</th>
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<td>5.00</td>
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<td>ab</td>
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<td>23.06</td>
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<td>15.88</td>
<td>21.13</td>
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<td>di</td>
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<tr>
<td>fo</td>
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<td>8.36</td>
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</table>

### Modes

- **Olivine**: Tr.
- **Pyroxene**: 34
- **Plagioclase**: 41
- **Opaques**: 8
- **Glass**: 16
- **Apatite**: 1

---

**DIABASE SEGREGATIONS**

Segregations consisting principally of medium-grained clinopyroxene and plagioclase have been found in hole 3, at depths below 32 feet (fig. 9). Two have been observed. The uppermost and largest segregation occurs between 32.8 and 34.1 feet (sample KI-228) and the smaller one, which is less than 0.1 foot thick, occurs at 38 feet. In thin sections of the largest segregation the equigranular clinopyroxene and plagioclase crystals form a hypidiomorphic to subophitic intergrowth with minor opaque minerals, olivine, apatite(?), and as much as 16 percent interstitial glass (fig. 13). No hy-

---

**Figure 13.—Photomicrograph of diabase segregation with plain light (A, upper) and crossed nicols (B, lower) showing hypidiomorphic-granular to subophitic clinopyroxene-plagioclase intergrowth. Large dark areas in upper middle and right middle parts are vesicles. Interstitial glass with needle-like apatite(?) crystals occurs around the long opaque crystal aggregate and in the left middle part.**
persthene was observed. The crystals commonly project into and also grow within the abundant irregular vesicles and form a local diktytaxiticlike texture.

Plagioclase, the most abundant mineral of the segregations (41 mode percent), occurs as subhedral laths and plates averaging 1 mm in length and 0.2 mm in width. The crystals are zoned ranging from An45 to An48, noticeably more sodic than the plagioclase in the olivine basalts. Subhedral to subophitic clinopyroxene, in subordinate amount (34 mode percent), occurs as unzoned and optically homogeneous crystals (2V=45°, γ=1.72) averaging 1 mm in diameter. The clinopyroxene has a composition of Wo25.5 En46.5 Fs38.0 (table 6), containing a somewhat greater proportion of ferrosilite than the groundmass augites of the olivine basalts. Olivine was not observed in thin section but was present in trace amounts in the heavy-mineral separates from the segregation. The olivine has an average composition of Fa42; slight zoning is apparent in some of the crystals. Although the segregation olivine is considerably richer in iron than the olivine in the basalt, the average magnesia content of the segregation olivine is still greater than that of the coexisting clinopyroxene. This relation supposedly indicates nonequilibrium (Muir and Tilley, 1957), a condition which appears untenable in view of the observed physical and mineralogical features in the segregation and enclosing rock. The opaque minerals are all strongly ferromagnetic and occur as elongate and equant crystals and as chains of eucrypt crystals.

**Table 6**—Composition of clinopyroxene from diabase segregation, sample KI-128A, hole 3

<table>
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<th>Chemical analysis</th>
<th>K2O</th>
<th>Na2O</th>
<th>CaO</th>
<th>MgO</th>
<th>Fe2O3</th>
<th>FeO</th>
<th>MnO</th>
<th>Al2O3</th>
<th>SiO2</th>
<th>WO3</th>
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<td>SiO2</td>
<td>51.04</td>
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<td>1.44</td>
<td>99.92</td>
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<tr>
<td>WO3</td>
<td>35.5</td>
<td>46.5</td>
<td>18.0</td>
<td>44.4</td>
<td>80.8</td>
<td>1.14</td>
<td>1.44</td>
<td>99.92</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

May contain some P2O5.

The interstitial glass (n=1.508) in the segregations represents an extreme tholeitic differentiate. Unlike the glass in the transient zone of crystallization, however, it was apparently solid when collected (fig. 12) and hence not in equilibrium with the crystal phases of the segregation at the collection temperature. In thin section the glass is colorless to very pale brown and contains fairly abundant needlelike growths of apatite (1). The work of Stewart (1962) indicates that the glass probably contains more than 60 percent SiO2.

A chemical and modal analysis of the diabase segregation is presented in table 5. The relatively low-bulk-silica content (50.32 percent), despite the predominance of silica-rich constituents (plagioclase, 55 percent SiO2; clinopyroxene, 51.04 percent SiO2; glass, 60 to 68 percent SiO2), is apparently due to the 8 percent of opaque minerals. An interesting, but inexplicable feature of the segregation not observed in other siliceous tholeiitic differentiates from Hawaii, is the extremely rich TiO2 content of 4.40 percent, which results in a high ilmenite norm of 8.36 percent. Moreover, in the ooze from hole 1, which is even richer in silica, the titania content is relatively low and comparable to titania values in other rocks of similar silica content. No abnormal changes in the remaining oxides, other than those attributable to fractional crystallization, are apparent.

**MELT**

The melt is that part of the lava lake with temperatures above 1065° C and is a viscous liquid containing variable amounts of crystalline olivine, minor clinopyroxene, and a trace or no plagioclase. Although the mineralogy and bulk chemistry of the melt (where sampled), crust, and original lavas are grossly similar, there is a marked enrichment in alkali content of the melt which appears to have significant petrologic implications. The melt in hole 1 (sample KI-112) is also abnormally high in titania, probably indicating some contribution of titaniferous interstitial liquid from the transient zone of crystallization.

In figure 14 are plotted the K2O:Na2O ratios, as a function of weight percent SiO2, for all the analyzed samples from the Kilauea Iki lava lake. Also plotted is the average composition of the 1959 lavas (K. J. Murata, unpub. data) that filled the lake. The outstanding features shown by this plot are: (1) the uniform K2O:Na2O ratios for most of the crust rocks, including those of the average 1959 lava, and (2) the anomalously high ratios for the two analyzed melt samples (KI-112, table 1; KI-168, table 2). Alkali ratios for the holocrystalline olivine basalts, most of the vitric-rich basalt in the transient zone of crystallization, and the average 1959 lava, consistently fall between 0.22 and 0.25, regardless of bulk-silica content. The melt in hole 1 (sample KI-112, depth 22.5 ft) and hole 2 (KI-168, depth 29.8 ft), on the other hand, have alkali ratios of 0.29 and 0.47 respectively. One sample, from the transient zone of crystallization (KI-165, depth 27.9 to 29.1 ft) about 1 foot above the melt in hole 2, also shows a marked increase in the alkali ratio. However, other samples from this zone (KI-164, depth 27.5 to 27.9 ft; KI-184, depth 31.8 to 32.8 ft; and KI-186, depth 34.3 to 35 ft) have a normal alkali con-
ALTERATION

The rocks of the crust show only minor affects from alteration. The most obvious alteration phenomenon is along the many cracks and under the small tumuli on the lake where the lava has acquired a reddish-brown hue due to a surficial oxidation of the iron by ascending hot vapors. Around the hotter and hence deeper cracks, the lavas are partially digested, leaving a rime of gelatinous silica, often in association with crystalline sublimates of sulfur and the anhydrous alkali sulfates, thenardite, and aphthitalite.

Within the crust some of the olivines undergo an incipient alteration consisting of an orange-brown stain around the margins of the crystals and along fractures penetrating the crystals. This staining first becomes apparent at a depth of about 2 feet below the surface and continues, generally without interruption, to about 12 feet above the crust-melt contact.

Chemically, the chief subsolidus change which occurs in the crust is the oxidation of iron. This change is graphically shown in figure 15, where the $Fe_2O_3$: $FeO$ ratios of the drill-hole samples are plotted against the depth at which they were collected. Although the points show scatter owing to the varying amounts of oxidation associated with cracks, the ratio indicates a decreasing degree of oxidation of iron with depth in the olivine basalt of the lake crust. The minimum value of $Fe_2O_3$: $FeO$ of about 0.1 at a depth of about 30 feet in hole 2 indicates the probable state of oxidation of juvenile or at least very recently...
erupted Kilauea lava. The increase of the ratio toward the top of the crust suggests that oxidation is proceeding from the top down and that the oxidizing agents are of atmospheric origin.

In figure 15 the offset of the average curve of points of hole 1 as compared with that of hole 2 is a measure of the oxidation which has taken place in the crust during the approximate 1-year interval between drilling and sampling of the two holes.

The measured increase in the oxidation state of the iron is probably the result of several processes. The late crystallization of opaque minerals at the base of the crust has already been noted (fig. 11). These opaques, which probably are chiefly magnetite, presumably grow at the expense of FeO in the glass (or interstitial melt). Further oxidation of iron in olivine occurs at rather shallow depth and at low temperature and produces the orange-brown staining referred to above.

DIFERENTIATION

Magmatic differentiation played and continues to play an important role in the crystallization of the tholeiitic rocks in the crust of the Kilauea Iki lava lake. During the 3-year period between December 1959 and December 1962, the crust has grown from 0 to more than 42 feet thick and fractional crystallization has resulted in compositional changes between 45.61 and at least 54.08 percent SiO₂. The petrographic and chemical data also strongly suggest that gas transfer may play an important role in desilication of the melt and that a process of transient alkali enrichment occurs in the melt immediately below the crust-melt contact. These changes in the composition of the crust and underlying melt have, without much doubt, occurred after the beginning of the formation of the crust and hence are not inherited, except indirectly, from original compositional differences in the lava of the 17 eruptive phases that filled the lake.

FRACTIONAL CRYSTALLIZATION

The changes in composition exhibited by most of the rocks are due to two processes of fractional crystallization: gravity settling of heavy crystals and filter pressing. Gravity settling has influenced the composition of the entire crust, but is more apparent—at the present time—only in the upper section. Filter pressing, however, is a more deep-seated process, dependent essentially on slow cooling, and is restricted to depths below about 20 feet.

In the upper 14 feet of crust the sinking of relatively heavy olivine crystals has produced a number of olivine-rich layers that contain as much as 37-mode-percent olivine and simultaneously leave a superstratum relatively poor in olivine (figs. 7, 8, and 9). The olivine accumulates rest upon thin layers of dense basalt, which probably represent partially digested portions of old crust emplaced during the period of crustal foundering at the end of the eruption. At depths of more than 14 feet there is a gradual but erratic decrease in olivine content (figs. 8 and 9). Olivine from this zone and the melt below is presumably building up a thick crystal accumulate at the base of the melt in the bottom of the lake.

The range in silica content from 45.61 to 49.61 percent in the crustal olivine basalts can be explained solely by the addition or removal of magnesian olivine by gravity settling (fig. 10). No evidence to support clinopyroxene removal by gravity settling, as advocated by MacDonald and Katsura (1961) has been found. On the basis of observations made during the 1959 summit eruption and 1960 flank eruption, augitic clinopyroxene begins to crystallize at about 1100° C (D. H. Richter, unpub. data). Hence, clinopyroxene can crystallize only in a zone extending upward from about 3 feet below the crust-melt contact (see temp profile, hole 2, table 1).

However, inasmuch as the crystals formed are small (max size 0.08 mm) and the melt is viscous, it appears unlikely that the rate of crystal settling is greater than the rate of crustal growth. Possibly, in time, with a thicker crust and less steep thermal gradient, the sinking of early-formed clinopyroxene may play a significant role in the differentiation of the lake lavas.

The efficacy of filter pressing as a differentiation mechanism in the lava lake is manifest by the presence of diabase segregations in hole 3 and the ooze from the bottom of hole 1. Both of these differentiates represent an interstitial liquid that was originally in equilibrium with crystallizing clinopyroxene and plagioclase in the transient zone of crystallization. Filter pressing becomes an effective separation process when a crystal mesh of sufficient cross-sectional density screens out crystals and allows only residual fluid to pass through. In the Hawaiian tholeiites, the Kilauea Iki lavas in particular, this condition is reached only after the crystallization of most of the magnesian olivine and some of the clinopyroxene and plagioclase. The expelled liquid was apparently injected into local dilatant zones, such as the continuously forming shrinkage cracks, at somewhat higher levels in the crust than the source of the liquid. The ooze in the drill hole, however, was artificially released by the drilling and thus allowed the interstitial liquid to exude from the zone of crystallization and to collect in the bottom of the hole.

The course of differentiation exhibited by the segregations and ooze is towards enrichment in K₂O, Na₂O,
and TiO₂ with increasing silica content (fig. 10). Alumina and total iron oxides remain virtually constant, and magnesia and lime show a marked decrease. Assuming that the liquid in the parent liquid-crystal melt (50 percent SiO₂) contained more than about 52 percent SiO₂, this trend would fall within a continuously changing clinopyroxene-plagioclase control field (Richter and others, 1964). The composition of the differentiates may also be affected by enrichment in potash and silica that is in addition to simple concentration through fractional crystallization.

A comparison of refractive indices (fig. 12) indicates that the ooze from hole 1 represents the residual liquid in an olivine basalt (5 to 10 percent modal olivine) after crystallization has been 76 to 78 percent complete. By inference from the index of refraction and the thermal-profile data in figure 12, this differentiate, which corresponds chemically to a quartz basalt, has a liquidus temperature of approximately 1040°C. The diabase segregations, because of their lower silica content, were evidently derived from less crystalline layers and at somewhat higher temperatures (1050°C to 1060°C) deeper in the zone of crystallization. Although residual liquids persist to levels where temperatures are as low as 970°C, the small volumes of these interstitially trapped liquids virtually preclude their egress in any substantial amount from the crust. However were it possible to continuously remove these lower temperature liquids, products approaching the composition of granite could readily be formed through fractional crystallization alone.

Analyses of the lava-lake rocks plotted on an AMF (alkali-magnesia-iron oxide) diagram (fig. 16) show only a very moderate iron enrichment, markedly different from the high iron trend of the Skaergaard (Wager and Deer, 1939) or the so-called Kilauea trend of Kuno and others (1957). The differentiation trend is practically identical with that shown by the Uwekahuna laccolith in Kilauea caldera (Murata and Richter, 1961) and is very similar to that of the Koolau basalts and intrusives on the Island of Oahu (Kuno and others, 1957). Kuno and his colleagues preferred the term “Kilauea trend” because of an iron-rich segregation vein in an olivine basalt flow in the wall of Kilauea caldera; however, as shown in figure 16, iron enrichment in Kilauea appears to be more the exception than the rule. Osborn (1959) has pointed out that changes in the partial pressure of oxygen in (pO₂) may be the controlling factor in determining which trend—iron enrichment or alkali enrichment—will be followed by a differentiating basaltic magma. However, it seems difficult to believe that in the similar environment of the flows, intrusives, and lava lake on Kilauea, changes in pO₂ would be of sufficient magnitude to affect the course of crystallization.

ALKALI TRANSFER

Petrochemical studies of melt samples from the lava lake strongly suggest that a zone enriched in alkalis, especially K₂O, occurs in the melt directly beneath the crust-melt contact. K₂O in this zone shows upward of 100 percent enrichment, resulting in a twofold increase in the K₂O:Na₂O ratio. After crystallization proceeds through this zone, the resulting holocrystalline rock shows a normal alkali content and alkali ratio. The thickness of the enriched layer is not known.

An attempt to explain this phenomenon is portrayed diagrammatically in the three sections in figure 17. At the end of the eruption in December 1959, the melt in the lake was probably nearly homogeneous, having an average K₂O:Na₂O ratio of 0.24 (fig. 17A). In fact, in 17 analyses of the lava that contributed to the filling of the lake the alkali ratio ranged only from 0.23 to 0.26 (K. J. Murata, unpub. data). In August 1960, when the crust was 22 feet thick, the sampled melt had a K₂O:Na₂O ratio of 0.29 (fig. 17B); and by April 1961 this ratio had increased to 0.47 (fig. 17C). The olivine basalt samples from the crust retain an average alkali ratio about the same as that of the original lavas, regardless of their depth in the crust. In the remaining great bulk of melt beneath the alkali-enriched zone, there is evidently a complimentary decrease in the alkali ratio; however, the actual values shown in figures 17B and C are highly conjectural. This process of alkali enrichment within the melt may be similar to the mechanism advanced by Kennedy (1955). He has
shown that under equilibrium conditions, water in a quiet magmatic melt will have a uniform partial pressure throughout its entire volume. To achieve this condition, water by means of diffusion will migrate to, and concentrate in, areas of least confining pressures and lowest temperatures. Utilizing the data of Morey and Hesselgesser (1952) who found an extremely high solubility of alkali silicates in superheated water vapor, Kennedy further reasoned that water and alkalis should tend to associate in a magma regardless of whether they are in solution or exist as a separate gas phase. Hence it appears very reasonable to expect that, in the lava lake, dissolved water in association with abundant alkalis will concentrate in the relatively low temperature-pressure zone beneath the crust-melt contact.

Upon crystallization of this enriched alkali-water zone and concomitant rapid decrease in pressure, the alkalis, in excess of that required to form plagioclase, are apparently released together with water vapor and other volatiles (fig. 17B, C). Migrating upwards, these volatile exhalations deposit crystalline sublimates of aphthitalite, thenardite, and sulfur in the cooler parts of the crust.

The petrologic implications of the alkali-enriched zone in the melt of the Kilauea Iki lava lake are great. Of particular importance is its bearing on the origin of the undersaturated alkali basalts. Macdonald (1949a, b) who has supported fractional crystallization to explain the genesis of the alkali basalts from a tholeiitic parent has also stressed the possible role played by alkali transfer (1949a, p. 1584; 1949b, p. 91). However, it was not until the first drilling in the lava lake that demonstrable evidence was found to sustain these views (Macdonald and Katsura, 1961). Additional data gathered subsequently from the lava lake and presented in this paper corroborates the presence of an alkali-rich melt, at least in part, and demonstrates that Macdonald's one sample was not exceptional.

If processes similar to those presently active within the lava lake occur in the shallow (1 to 5 km deep) magma chambers of the Hawaiian volcanoes, there is convincing argument to support the derivation of the alkali basalts from a primary tholeiitic magma. Throughout the early stages of Hawaiian volcanism, tholeiitic lavas are erupted frequently and voluminously without undergoing prolonged storage in the volcanic pile. With decadency, however, the tempo of eruptive activity declines. The processes of alkali enrichment and silica transfer in the Kilauea Iki lava lake may have upwards of hundreds of years to operate rather than only a few, and the lavas shift to the alkalic type. It is also significant that the total volume of alkalic lava erupted is very minor compared with the tremendous volume of tholeiitic basalt composing the main bulk of the volcano. During the long periods of quiescence in the waning period of volcanism, extremely slow cooling and relatively high lithostatic load should enhance the concentration of alkalis in the upper levels of the stagnant magma chambers and may also further the process of silica transfer. Although desilication of the magma through gas transfer still evades quantitative evaluation, it is an apparently important process and may well be the key in explaining the undersaturation of the alkalic basalts. Slow cooling will also significantly increase the efficacy of fractional crystallization in the magma. Not only will magnesian olivine be removed from the upper part of the melt through gravity settling, but the early-crystallizing high-silica augitic clinopyroxenes should settle out as well. The formation of magnesian olivine and augitic accumulates in the shallow magma chambers is indicated by the great

![Diagram](image-url)

**Figure 17**—Hypothetical sections through Kilauea Iki lava lake showing mechanism of alkali enrichment. Thickness of crust is exaggerated and solidified base is omitted.
abundance of periodotite and pyroxenite nodules in the alkalic flows (Richter and Murata, 1961), whereas such inclusions are almost unknown in Hawaiian tholeiites.

Eruptions of the postulated alkali-rich melt could be effected by an increase in vapor pressure that accompanies the concentration of H2O in the residual melt in the uppermost parts of the magma chambers. The need for additional primary material from the mantle to act as an accessory force to cause eruption, as in the tholeiitic eruptions, does not appear necessary. An explosive release of water vapor would follow any fracturing of the magma chamber. Indeed, the scattered occurrence of vents and apparent explosive character of Hawaiian alkalic eruptions implies a fairly shallow origin for the magma, such that fracturing from the magma chamber to the surface is geologically feasible.

Recently, evidence has been found to indicate the possible beginning of a shift to alkalic tendencies in Kilauea Volcano (J. G. Moore and D. H. Richter, unpublished data). The K2O: Na2O ratios of prehistoric and early historic tholeiitic basalts from Kilauea average about 0.19, while recent basalts average 0.24. This change is in the same direction as that of the change from tholeiitic to alkalic basalt and also of that in the changes observed in the lava lake.

Thus, it appears plausible that a combination of alkali and silica transfer and fractional crystallization of clinopyroxene within shallow magma chambers may well explain the change from tholeiitic to alkalic basalt observed in Hawaiian volcanoes. This is opposed to the views of Yoder and Tilley (1962) who, principally on the basis of laboratory studies, believe that neither a tholeiitic nor alkalic magma is the parent of the other but that each is derived as separate fluids from the mantle. Although our arguments have been largely qualitative, further investigations on the slowly crystallizing Kilauea Iki lava lake should significantly add to the presently meager knowledge of magmatic chemical-physical processes.

REFERENCES


