Hydrothermal Alteration in Research Drill Hole Y-3, Lower Geyser Basin, Yellowstone National Park, Wyoming

U.S. GEOLOGICAL SURVEY PROFESSIONAL PAPER 1054-C
Hydrothermal Alteration in Research Drill Hole Y-3, Lower Geyser Basin, Yellowstone National Park, Wyoming

By KEITH E. BARGAR and MELVIN H. BEESON

HYDROTHERMAL STUDIES IN YELLOWSTONE NATIONAL PARK, WYOMING

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HYDROTHERMAL STUDIES IN YELLOWSTONE NATIONAL PARK, WYOMING

HYDROTHERMAL ALTERATION IN RESEARCH DRILL HOLE Y-3,
LOWER GEYSER BASIN, YELLOWSTONE NATIONAL PARK, WYOMING

By Keith E. Bargar and Melvin H. Beeson

INTRODUCTION

Research diamond-drill hole Y-3 is one of 13 holes completed by the U.S. Geological Survey at selected sites in hot-spring and geyser areas of Yellowstone National Park (fig. 1). The holes were drilled in 1967 and 1968 in order to obtain detailed physical and chemical data on the shallow part of a high-temperature geothermal system (White and others, 1975). Drill hole Y-3 is located at the western margin of the Pocket Basin hydrothermal explosion crater debris ridge (Muffler and others, 1971). The drill-hole surface occurs at an elevation of ~2,193 m; Y-3 is about 76 m north of and 2 m higher than Ojo Caliente Hot Spring, a major thermal feature in the area (White and others, 1975).

Drilling of Y-3 (using water as the circulating fluid) began on June 8, 1967, and was terminated on June 28, 1967, after reaching a depth of 156.7 m. The drill hole was later plugged with cement in 1969. The core diameter is 10.2 cm for the interval from the surface to 9.8 m, 6.0 cm for the interval from 9.8 to 27.4 m, and 4.4 cm for the interval from 27.4 m to the drill-hole bottom. Core recovery, accomplished by wire-line method, averaged 65 percent from the surface to 28.7 m, only 3 percent from 28.5 to 41.1 m, and 94 percent in the remainder of the drill hole.

The drill core, which was usually obtained in 1.5- or 3-m intervals, was logged at the drill site by R.O. Fournier, A.H. Truesdell, and D.E. White as soon as possible following core recovery. At that time approximately 370 pieces of drill core were selected for detailed laboratory study on the basis of being representative of an interval or because of secondary mineral deposits in veins, vugs, or fractures. This “skeleton” core was systematically studied by optical (44 thin sections and polished thin sections), X-ray diffract (more than 700 X-ray diffractograms), electron microscope, and scanning electron microscope (SEM) methods. Whole-rock chemical analyses by rapid rock methods and semiquantitative spectrographic analyses for 16 selected Y-3 core samples are reported in Beeson and Bargar (1984). Bulk-rock and powdered-rock density measurements were also obtained for these 16 core samples. Two of the more abundant hydrothermal minerals (chlorite and laumontite) were concentrated by heavy liquids for conventional rock analysis.

Physical data for the 13 Yellowstone research drill holes are reported in White and others (1975). The highest temperature recorded in Y-3 was 196.0 °C at a depth of ~150.6 m (fig. 2), and the maximum near-bottom fluid pressure was ~1,497 kPa.

Bargar and others (1973) indicate that the Y-3 drill hole encountered at least two major water-producing zones at depths of 28 and 88 m, and they provide a chemical analysis of the slightly alkaline water from the 88-m aquifer (table 1, column 2). More recently a chemical analysis of water from the bottom of the Y-3 drill hole was reported by Keith and others (1983), and the analysis is reproduced here in table 1, column 1. This water is slightly more concentrated in the major constituents (Na, HCO₃, and Cl) than water from the 88-m aquifer. Silica was not reported in either of the two analyses; however, SiO₂ (along with Na, HCO₃, and Cl) is one of the four main components of nearby Ojo Caliente Hot Spring water (table 1, column 3) (Thompson and Yadav, 1979).
HYDROTHERMAL ALTERATION IN RESEARCH DRILL HOLE Y-3, WYOMING

ACKNOWLEDGMENTS
The authors would like to thank T.E.C. Keith and D.E. White for their critical reviews and helpful discussion of the manuscript. R.L. Oscarson assisted in obtaining SEM micrographs, and L.J.P. Muffler provided us with his notes on the petrography of some of the Y-3 drill core.

STRATIGRAPHY
SURFICIAL DEPOSITS
The upper 1.6 m of Y-3 drill core consists of buff-to-white clayey (smectite and kaolinite) alluvial soil that contains some sand-sized grains of black spherical obsidian and clear botryoidal opal and angular-to-subrounded light-gray rhyolite pebbles.

Between 1.6 and 42.2 m the drill core is composed of red, brown, gray, or green hydrothermally cemented siltstone, sandstone, and conglomerate. The massive to thinly bedded sediments have dips that range from horizontal (perpendicular to drill core that is presumed to be vertical) to about 30°; several samples display crossbedding or contorted bedding. The conglomeratic material is poorly sorted and contains mostly rhyolite and obsidian clasts, although a few samples have sandstone pebbles. These sediments were deposited during the Pinedale glacial episode (≥45,000–14,000 years B.P.)
Figure 2.—Plot showing depth distribution of unaltered obsidian, α-cristobalite, and hydrothermal alteration minerals from drill core Y-3. Left column shows a generalized stratigraphic section of rock units penetrated by the drill hole. The unnamed rhyolite flow and the Nez Perce Creek rhyolite flow are included in the Central Plateau Member of the Plateau Rhyolite. Horizontal lines adjacent to left column represent distribution of samples studied. Only a few core fragments and drill cuttings were recovered between 28.5 and 41.1 m, and the distribution of hydrothermal minerals in the interval, as suggested by these few samples, is shown by vertical dotted lines. Width of mineral columns given as an estimate of relative abundance based on X-ray diffraction analyses and microscopic observations. Vertical continuity in mineral presence between samples is assumed except where a mineral abundance becomes zero, in which case the zero point is arbitrarily placed 0.3 m from the last occurrence of the mineral. Dashed curve connects bottom-hole temperature measurements obtained as drilling progressed. Solid curve shows reference boiling temperature for pure water assuming a surface altitude of ~2,329 m and a water table at the surface. Temperature data from White and others (1975).
(Muffler and others, 1971; Waldrop and Pierce, 1975; Pierce and others, 1976; Muffler and others, 1982). Hydrothermal alteration is extensive in samples recovered throughout most of the interval, and hydrothermal minerals include opal, β-cristobalite, quartz, chaledony, analcime, clinoptilolite, mordenite, calcite, fluorite, allophane, halloysite, kaolinite, chlorite, smectite, lepidolite, aegirine, pektolite, gypsum, alunite, pyrite, and hematite.

**VOLCANIC ROCKS**

From 42.2 to 71.2 m the drill core penetrated the Nez Perce Creek rhyolite flow of the Central Plateau Member of the Pleistocene Plateau Rhyolite (Christiansen and Blank, 1974), which was extruded about 150,000 years ago (Waldrop and Pierce, 1975; White and others, 1975; Leeman and Phelps, 1981). Core samples from the Nez Perce Creek flow frequently consist of bleached rhyolite flow or flow breccia that is extensively fractured and in some places is quite vesicular. Predominant magmatic minerals are quartz and sanidine; pyroxene, plagioclase, and magnetite are minor constituents. Evidence for vapor-phase mineralization is sparse, although a few cavities contain hydrothermal quartz druses that appear to be pseudomorphs after vapor-phase alkali feldspar crystals. Spherulitic textures are common throughout the rhyolite flow, but α-cristobalite, which frequently forms due to devitrification of cooling rhyolite flows, is rare and apparently has been recrystallized to quartz. Hydrothermal quartz and chaledony are abundant on fractures and as vesicle fillings throughout much of this core interval. Other open-space hydrothermal minerals identified in the Nez Perce Creek flow include mordenite, dachiardite, aegirine, pektolite, calcite, fluorite, smectite, illite-smectite, chlorite, kaolinite, hematite, pyrite, and a 10-Å mica(?).

Below 71.2 m is an older, unnamed rhyolite flow that apparently is not exposed anywhere at the surface within the Yellowstone Caldera. This flow is also correlated with the Central Plateau Member of the Pleistocene Plateau Rhyolite by Christiansen and Blank (1974). The unnamed rhyolite flow is probably not much older than the Nez Perce Creek flow because all of the lava flows of the Central Plateau Member were extruded about 200,000–70,000 years ago (Christiansen and Blank, 1972). In drill core Y-3, this unnamed rhyolite flow displays conspicuous flow banding and vesicularity. Quartz and sanidine are the most abundant magmatic minerals, and pyroxene, plagioclase, and magnetite are minor primary constituents. No vapor-phase or devitrification minerals were identified, although parts of the drill core have a spherulitic texture. The core is extensively fractured, and hydrothermal minerals are quite abundant as vesicle and fracture fillings, to the extent that many of the fractures are completely self-sealed. Hydrothermal minerals found in this partially bleached rhyolite flow include quartz, chaledony, dachiardite, laumontite, mordenite, stilbite, yugawaralite, truscottite, calcite, fluorite, smectite, illite-smectite, illite, chlorite, pyrite, and hematite.

**HYDROTHERMAL ALTERATION MINERALOGY**

Figure 2 shows the distribution of unaltered obsidian, devitrification α-cristobalite, and hydrothermal minerals (opal, β-cristobalite, quartz, chaledony, analcime, clinoptilolite, dachiardite, laumontite, mordenite, calcite, fluorite, kaolinite, halloysite, chlorite, smectite, illite-smectite, lepidolite, aegirine, pektolite, gypsum, pyrite, and hematite) with depth in the Y-3 drill core. Vapor-phase, magmatic, and a few hydrothermal minerals with only minor distribution in the drill core (stilbite, yugawaralite, allophane, truscottite, and alunite) are not shown in the figure.

The only unaltered obsidian occurs in the coarse-grained sediments of the upper 10 m of the drill core.
Below that depth, clastic obsidian grains are altered to clinoptilolite and smectite.

Silica minerals

Opal and β-cristobalite

Sediments in the upper 14.0 m of the drill core are mostly cemented by clear-to-grayish opal. Opal also occurs in a few veins and vugs where the mineral usually has a botryoidal habit; however, one cavity at 4.7 m is partly filled by brown banded opal that grades upward to clear hyalite. X-ray diffraction analyses of two samples at 10.8 and 18.9 m show the presence of β-cristobalite associated with opal cement in the shallower deposit and a chalcedony vein in the deeper sample.

Quartz and chalcedony

Red-to-brown, green, and white-to-clear chalcedony occurs throughout most of the Y-3 drill core below 18.9 m. The cryptocrystalline silica was deposited as botryoidal, irregular-to-massive, or horizontally layered cavity and fracture fillings. In addition, chalcedony forms the cementing material for a few pieces of conglomerate or brecciated drill core. It also appears to be the earliest hydrothermal mineral deposited in the Y-3 drill core, although some early chalcedony contains disseminated, minute pyrite crystals. A few chalcedony cavity fillings display microscopic convoluted bedding. Similar sedimentary structures in chalcedony cavity and fracture fillings of drill core Y-5 were interpreted by Keith and Muffler (1978) as being deposits of amorphous silica floc (opal) that later recrystallized to chalcedony.

Tiny (0.025–0.5 mm), isotropic, euhedral, trapezohedral analcime crystals (fig. 3) occur in the groundmass and as vesicle fillings in the Y-3 drill core from 14.2 to 15.3 m, 22.8 to 24.1 m, and 27.2 to 28.5 m. Quartz is closely associated with analcime and appears to be pseudomorphous after groundmass analcime at 23.8 m. Six electron microprobe chemical analyses of analcime from 27.5 m (table 2) show that the mineral is a pure analcime (sodium-rich) end member. The Y-3 analcime analyses have very high silica-to-alumina ratios (2.4–3.1) for analcime and are similar both to a microprobe analysis of analcime reported for drill core Y-8 (Keith and others, 1978b) and to conventional rock analyses of two analcime separates from drill core Y-1 (M.H. Beeson, unpub. data, 1976). Other Yellowstone drill cores that contain analcime are Y-2 and Y-13. In drill core Y-2 the mineral appears to be a sodium-rich wairakite (Bargar and Beeson, 1981), and in drill core Y-13 both analcime and wairakite appear to be present in the same sample (Bargar and others, 1981). Drill cores Y-1 and Y-8 mostly contain Na- and K-rich hydrothermal minerals and do not contain Ca-rich minerals (except for local bladed calcite) (Honda and Muffler, 1970; Keith and others, 1978b). Conversely, drill cores Y-2 and Y-13 contain several Ca-rich hydrothermal minerals (Keith and others, 1978a; Bargar and Beeson, 1981).
TABLE 2. - Chemical composition of some zeolite minerals from Y-3 drill core
[All analyses except for laumontite (No. 5) and H₂O⁺ and H₂O⁻ of mordenite were determined by electron microprobe]

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<td>Cat. bal. error (%)</td>
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1Sample numbers correspond to depth in meters.
3H<sub>2</sub>O<sup>+</sup> and H<sub>2</sub>O<sup>-</sup> determined by conventional rock-analysis method. Analyst: S. T. Neil.
4Average of five analyses.
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Numbers of atoms on the basis of 48 oxygens

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| Al | 5.05 | 4.97 | 4.00 | 4.23 | 4.53 | 4.45 | 4.27 | 4.80 | 4.33 | 4.63 | 5.27 | 5.12 |
| Fe | 0.00 | 0.00 | 0.00 | 0.01 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 |
| Mg | 0.01 | 0.00 | 0.00 | 0.01 | 0.01 | 0.01 | 0.00 | 0.00 | 0.00 | 0.01 | 0.01 | 0.00 |
| Mn | 0.02 | 0.01 | 0.00 | 0.01 | 0.00 | 0.00 | 0.00 | 0.00 | 0.01 | 0.00 | 0.02 | 0.01 |
| Ca | 1.89 | 1.92 | 1.51 | 1.56 | 1.60 | 1.54 | 1.30 | 1.74 | 1.53 | 1.69 | 2.07 | 1.99 |
| Na | 0.55 | 0.53 | 0.77 | 0.58 | 0.66 | 0.76 | 0.66 | 0.86 | 0.75 | 0.67 | 0.57 | 0.69 |
| K | 2.24 | 2.25 | 3.35 | 3.35 | 3.7 | 3.46 | 3.7 | 4.0 | 2.75 | 2.74 | 2.33 | 2.38 |
| Si/Al | 3.77 | 3.85 | 4.99 | 4.68 | 4.31 | 4.39 | 4.65 | 4.00 | 4.55 | 4.20 | 3.57 | 3.70 |
| Cat. bal. | 3.97 | 3.27 | 3.66 | 3.25 | 3.93 | 3.39 | 3.49 | 3.67 | 3.85 | 3.56 | 3.25 | 3.87 |
| error (%) | 8.7 | 6.7 | -3.7 | 4.0 | 6.8 | -4 | 8.4 | 1.9 | 2.7 | 7.6 | 6.3 | 3.5 | -10.9 | 10.2 | -2.4 | 9.3 | 4.9 |</p>
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Numbers of atoms on the basis of 48 oxygens

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1Sample numbers correspond to depth in meters.
2Average of two analyses.
3Average of five analyses.
Clinoptilolite

Clinoptilolite, usually associated with devitrification α-cristobalite in drill core Y-3 (fig. 2), was identified by X-ray diffraction analyses as the cementing material in finer grained sediments at 7.3–14.2 m. Clinoptilolite is the only zeolite mineral in this interval. The nearly isotropic mineral also occurs as cement in finer grained siltstone and sandstone and was deposited as tiny (0.005–0.012 mm), clear, tabular crystals that line cavities (fig. 4) in conglomerate between 16.2 and 26.3 m. X-ray diffraction traces of 12 samples from these two zones showed no change in position or intensity of the (020) X-ray peak at about 9 Å after heating according to the method of Mumpton (1960), and the mineral was identified as clinoptilolite.

A single microprobe chemical analysis of clinoptilolite was obtained from 12.4 m (table 2). This analysis indicates that clinoptilolite at this depth is low in silica, contains very little calcium, and is rich in sodium and, in particular, potassium. The Y-3 clinoptilolite is much lower in calcium than most other heulandite-group zeolites from Yellowstone drill cores, and it is nearly as rich in potassium as some clinoptilolites from drill cores Y-7 and Y-8 (Keith and others, 1978b) (fig. 5).

Three core samples from 24.2 to 26.3 m and drill cuttings recovered from about 29–30 m are characterized by splitting of the (020) X-ray peak into three peaks at about 8.9, 8.7, and 8.3 Å following heat treatment. No chemical analyses were obtained for the three core samples; however, we would predict that they contain more calcium than the above clinoptilolite samples and are probably intermediate heulandites (Alietti, 1972; Boles, 1972).

Dachiardite

Clear dachiardite crystals are widely scattered between 60.2 and 80.9 m in the Y-3 drill core. The rare, radiating, blocky crystals are found deposited in cavities along with earlier quartz, later mordenite, and smectite (figs. 6 and 7), and they are later than chalcedony in a few vein fillings.

Microprobe chemical analyses of dachiardite crystals from 60.4 and 73.2 m in drill core Y-3 are given in table 3. A ternary plot of the exchangeable cations (Ca-Na-K) content in Y-3 dachiardite compared with other published dachiardite analyses is shown in figure 8. The published analyses are divided into two groups: (1) sodium-rich dachiardite, and (2) intermediate dachiardite (Alberti, 1975; Wise and Tschernich, 1978; Bonardi, 1979; Bonardi and others, 1981). A drill core Y-2 dachiardite analysis (Bargar and Beeson, 1981) plots near the latter group and several dachiardite analyses from sample Y3-60.4 plot near the intermediate group, but these crystals are more calcium rich. Chemical analyses of water from the Y-3 drill hole (table 1) suggest that calcium is lost (precipitates) as the water ascends to the surface (Bargar and others, 1973). The Y3-73.2 sample is very calcium rich, and, to the best of our knowledge, this

![FIGURE 4.—Scanning electron micrograph of intergrown tabular clinoptilolite crystals from 14.0 m in drill core Y-3 (after Bargar and others, 1981).](image)

![FIGURE 5.—Ca-Na-K ternary diagram for heulandite-group zeolites from Yellowstone drill cores. Data sources as follows: C-1 (Fenner, 1936), V-1 (Honda and Muffler, 1970), Y-2 (Bargar and Beeson, 1981), Y-7 and Y-8 (Keith and others, 1978b), Y-6 (Bargar and Beeson, 1984), Y-3 (table 3, this report).](image)
sample is the only naturally occurring Ca-rich dachiardite ever reported. Ca-rich dachiardite has been produced in the laboratory using devitrified rhyolite as the starting material (Knauss and Beiriger, 1984).

LAUMONTITE

Many fractures and cavities below 73.2 m in the Y-3 drill core contain white prismatic laumontite crystals that are up to about 1 mm in length. SEM studies show that laumontite was deposited later than quartz and earlier than mixed-layer illite-smectite. Some laumontite crystals appear to have been subjected to partial leaching, and the illite-smectite was deposited in the resulting cavities (fig. 9).

Chemical analyses of laumontite from 73.2 and 126.9 m (by electron microprobe and conventional rock analysis methods) are given in table 2. The Y-3 laumontite is calcium rich and is very similar to several microprobe analyses of hydrothermal, burial metamorphic, and surficial hot-spring-deposited laumontite (McCulloh and others, 1981). The measured temperature range at which Y-3 laumontite was found was about 167–196 °C.

MORDENITE

Mordenite is the most abundant zeolite mineral found in the Y-3 drill core. Scattered mordenite deposits occur in many vugs and fractures throughout much of the drill core as radiating, long, thin, fibrous crystals (fig. 7), as

**FIGURE 7.—** Scanning electron micrograph of cavity filling from 60.2 m in the Y-3 drill core. Micrograph shows, in apparent depositional sequence, quartz crystals (upper left and lower right), dachiardite (blocky crystals extending diagonally from lower left to right center), radiating fibrous mordenite crystals, and numerous tiny smectite crystal clusters (after Bargar and others, 1981).

**FIGURE 8.—** Ca-Na-K ternary diagram for Y-3 dachiardite compared with other published dachiardite analyses. Data sources as follows: Y3-60.4 and Y3-73.2 (table 3, this report), Y2-20.7 (Bargar and Beeson, 1981), Pacific Northwest (Wise and Tschernich, 1978), Italy (Bonardi, 1979; Alberti, 1975), Canada (Bonardi and others, 1981).
mats of interwoven fibers, or as needlelike crystals (fig. 10). SEM studies show that mordenite formed later than silica and most other zeolite minerals and earlier than some clay deposits (figs. 7 and 10).

Electron microprobe chemical analyses of coarse mordenite fibers from 91.4 m are given in table 2 along with H$_2$O values that were determined by conventional rock analysis methods. None of the microprobe analyses total to 100 percent even with the accurate H$_2$O values, and cation balance errors are high for all but two (1 and 4 in table 2) of the analyses. Nevertheless, the microprobe analyses indicate that sodium and calcium are the major cations and that potassium is a minor constituent; these results are similar to that from a conventional rock analysis of mordenite by Sheppard and Gude (1969) (fig. 11).

YUGAWARALITE

Yugawaralite is a rare zeolite mineral that has been reported from only a few locations in the world (Zeng and Liou, 1982). Clear, tabular yugawaralite crystals (fig. 12) were deposited in vugs, veins, and fractures of four scattered samples between 71.8 and 75.2 m, where the temperature measured during drilling was about 170 °C. The open-space yugawaralite crystals were deposited later than chalcedony and quartz and earlier than clay. Other zeolite minerals (mordenite, dachiardite, laumontite, and stilbite) appear to be closely associated with yugawaralite; calcite, smectite, pyrite, and chlorite also occur in the yugawaralite-bearing samples.

The idealized unit cell content of yugawaralite is Ca$_4$(Al$_8$Si$_{20}$O$_{66}$)16H$_2$O (Barrer, 1982). Several microprobe chemical analyses of yugawaralite from the Y-3 drill core at 73.2 m (table 2) show that calcium is the dominant cation and that only minor amounts of sodium and potassium are present. The Y-3 chemical analyses are very similar to analyses of yugawaralite made by Eberlein and others (1971), Pongiluppi (1977), Wise (1978), and Zeng and Liou (1982).

STILBITE

Stilbite, a fairly common zeolite mineral in andesitic and basaltic volcanic terranes (Deer and others, 1963), is extremely rare in the rhyolitic rocks of the Yellowstone volcanic region. In fact, stilbite was only tentatively identified by X-ray diffraction from a single vein filling at 73.2 m (fig. 13) along with yugawaralite, dachiardite, laumontite, quartz, and chalcedony.

CALCITE

White, bladed calcite crystals (up to about 3 cm long) line cavities and fractures in the Y-3 drill core as shallow as 12.1 m. Calcite is sparsely distributed in the glacial sediments and the Nez Perce Creek rhyolite flow, but below about 100 m in the unnamed rhyolite flow, which composes the lower one-third of the drill core, calcite is present in most samples studied. Calcite appears to
have formed in the middle of the paragenetic sequence of minerals and is generally deposited later than chalcedony, pyrite, hematite, quartz, and chlorite, and before later quartz, zeolite, smectite, and illite-smectite.

**FLUORITE**

According to Browne (1978), fluorite, a common hydrothermal mineral, appears to be an uncommon hydrothermal deposit in presently active geothermal systems, and it has been reported only from Java and from several drill cores in Yellowstone National Park. Fluorite in drill core Y-3 is very sparse above about 91 m but is somewhat more abundant below that depth. White, spherical fluorite was identified in the lepidolite-aegirine-pectolite zone at about 27.5 m. Deeper fluorite deposits are usually massive or consist of clear octahedral crystals that formed earlier than clay and zeolite minerals and later than early quartz crystals. Fluorite is observed in association with calcite, but the order of deposition is not clear. A chemical analysis of water from 88 m, at the top of the lower fluorite zone, shows a high fluoride content (about 30 ppm) (Bargar and others, 1973).

**CLAY AND MICA MINERALS**

**KAOLINITE**

White to yellow-stained kaolinite was identified by X-ray diffraction in fractures, vugs, and whole-rock alteration in six zones dispersed between the surface and 71.3 m in drill core Y-3 (fig. 2). In the upper zone (1.4–5.8 m), kaolinite is found in association with opal, alunite, allophane, halloysite, pyrite, hematite, smectite, and gypsum. Much of this assemblage probably

**EXPLANATION**

Drill hole Depth (m)
• Y3-91.4
○ Sheppard and Gude (1969)

**Figure 11.**—Ca-Na-K ternary diagram for microprobe chemical analyses of mordenite from 91.4 m in drill core Y-3 compared with a mordenite analysis given by Sheppard and Gude (1969).

**Figure 12.**—Scanning electron micrograph of euhedral crystals of yugawaralite with later smectite deposits at 73 m in drill hole Y-3 (after Bargar and others, 1981).

**Figure 13.**—Scanning electron micrograph of blocky stilbite(?) crystals (St) intergrown with tabular yugawaralite (y) and later smectite (s) on a fracture surface at 73.2 m in drill core Y-3 (after Bargar and others, 1981).
formed as a result of acid-sulfate alteration. A similar mineral assemblage occurs in Yellowstone drill core Y-11 from a vapor-dominated geothermal system (Muffler and Bargan, 1974; Bargan and Muffler, 1982). Schoen and others (1974) also attributed a similar assemblage of minerals from Steamboat Springs, Nevada, to surficial acid (H$_2$SO$_4$) alteration. Two additional kaolinite zones (41.7–44.4 m and 68.7–71.3 m) contain associated smectite and pyrite and are nearly coincident with stratigraphic breaks between the glacial sediments and the top of the Nez Perce Creek rhyolite flow and between the Nez Perce Creek flow and the older, unnamed rhyolite flow. However, fossil zones of surficial acid alteration are precluded in these two zones because the kaolinite is a late hydrothermal product and is deposited on top of quartz. The kaolinite may have formed due to alteration of K-feldspar by descending meteoric waters that were acidified by oxidation of pyrite. Fenner (1936) describes a similar process for kaolinite formation in drill core C-II (fig. 1).

**HALLOYSITE AND ALLOPHANE**

At 1.7 m, kaolinite is the predominant clay mineral; however, some halloysite and allophane are also present. Halloysite has a broad X-ray diffraction peak at about 7.7 Å, which, upon heating to 140 °C for 15 hours, becomes sharper, higher, and shifts to about 7.3 Å (Birkeland, 1969). In addition, one of the X-ray traces of this sample contained a low, broad ~11.5 Å peak that was not affected by heating and might be an allophane peak; allophane is commonly associated with kaolinite and halloysite (Deer and others, 1962). Subsequent testing, involving a small amount of powdered sample placed upon phenolphthalein-treated filter paper to which a drop of 1M NaF was later added, confirmed the identification of allophane by turning the indicator paper to a pinkish-red color (Fieldes and Perrott, 1966).

**CHLORITE**

Chlorite occurs throughout most of the two rhyolite flows as fillings in fractures and cavities and as whole-rock alteration adjacent to a few fractures. Open-space chlorite deposits formed later than chalcedony, hematite, quartz, smectite, and mixed-layer illite-smectite, and before late quartz crystals, late pyrite, and zeolite minerals. X-ray diffraction analyses show that the (001) and (003) X-ray peaks are weak and that the (002) and (004) reflections are strong, which suggests that the Y-3 chlorite is an iron-rich mineral (Grim, 1968). The X-ray data were confirmed by a wet chemical analysis of chlorite scraped from a large cavity at 108.3 m (table 4). This analysis shows that the Y-3 chlorite is iron rich [Fe$^{2+}$/Fe$^{2+} + $Mg$^+$] = 0.914]. Very small amounts

| TABLE 4. -- Chemical composition of chlorite from 108.3 m in drill core Y-3, 1, 2 |
|-----------------|-----------------|
| Weight percent  |                |
| SiO$_2$         | 31.65           |
| TiO$_2$         | 1.15            |
| Al$_2$O$_3$     | 20.55           |
| FeO             | 28.11           |
| MnO             | 1.47            |
| MgO             | 1.49            |
| CaO             | 0.43            |
| Na$_2$O         | 0.44            |
| K$_2$O          | 1.64            |
| H$_2$O          | 9.07            |
| H$_2$O$^{-}$     | 0.74            |
| F               | 0.12            |
| Total           | 99.79           |
| O$_2$F          | 0.05            |

Numbers of atoms on the basis of 36 oxygens

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1X-ray diffraction shows about 1-2 percent quartz and hematite impurities.

(<–2 percent) of quartz and hematite contaminants were identified on an X-ray diffraction trace of the analyzed sample, and these could account for the somewhat high SiO$_2$ and Fe$_2$O$_3$ values. A low MgO value is not unexpected because of the low MgO content of both rocks and waters in drill hole Y-3, and a high MnO value is possible in chlorite (Deer and others, 1966); however, a high K$_2$O value is difficult to explain unless the sample was contaminated by phenocryst sanidine. No feldspar was detected by X-ray diffraction.
SMECTITE

White, light-gray, or green smectite is the predominant clay mineral in much of the upper 80.6 m of the Y-3 drill core. The smectite commonly occurs as an alteration product of glass, and it was deposited in cavities and on fractures. X-ray diffraction analyses show that the smectite basal (001) spacing varies from about 12 to 15 Å and expands to about 17 Å when treated with ethylene glycol for 1 hour at 60 °C. In the lower part of this range (12.5 Å) sodium is the exchangeable cation, and at the upper end of the range (14.5-15.5 Å) calcium should be the exchangeable cation (Grim, 1968). Using these criteria, calcium is the predominant cation in the smectite above 11.3 m and from 48.8 to 80.6 m in the Y-3 drill core; between these depths (11.3-48.8 m), sodium is the major cation, with occasional calcium-rich samples.

ILLITE-SMECTITE

Mixed-layer illite-smectite occurs as open-space fillings and whole-rock alteration below 80.6 m and can be found in a few shallower samples. X-ray diffraction traces show an air-dried basal (001) peak at about 10-12 Å for the mixed-layer clay that splits into two peaks at about 11-14 Å and 9.1-9.9 Å following glycolation. According to data given by Hower (1981), these samples appear to contain about 60-80 percent illite layers and are probably allevardite ordered.

LEPIDOLITE

Lepidolite, generally considered to be a high-temperature vein, pegmatite, granite, or aplite-associated deposit (Deer and others, 1966), was identified as an intergranular cavity filling in conglomerate and coarse-grained sandstone at 23.8 m and between 27.2 and 28.5 m in the Y-3 drill core, where the temperature was about 130-140 °C (Bargar and others, 1973). The polycrystalline lepidolite mats (fig. 14) appear to have been deposited later than analcime and quartz and earlier than pectolite and aegirine. Other hydrothermal minerals identified in the lepidolite-bearing samples include chalcedony, calcite, fluorite, kaolinite, and pyrite. X-ray diffraction and chemical data for the polylithionite (a lepidolite end member) are given in Bargar and others (1973). According to Bargar and others (1973), the water that deposited lepidolite in the Y-3 drill hole is anomalously rich in lithium and fluoride, which are the two chemical constituents necessary for the formation of lepidolite. A barely perceptible 10-Å mica X-ray diffraction peak was noted in a few scattered samples at 1.4, 13.0-15.2, 43.8-47.0, and 65.5-66.1 m. The 10-Å mica was not observed in thin section, and its origin is uncertain.

AEGIRINE

In drill core Y-3, bright-green, pleochroic aegirine occurs as spherical or radiating clusters of euhedral bladed-to-prismatic crystals that are up to about 15 µm in length (figs. 15A, B). Aegirine is closely associated with pectolite at depths of 26.3-28.5 and 65.5-66.3 m; in the shallower zone, the two minerals were deposited in intergranular cavities in coarse sediments, whereas in the deeper zone, the minerals were deposited in fractures and cavities formed by leaching of sanidine phenocrysts. Aegirine is a fairly rare mineral in modern geothermal systems; previously, it was identified in Yellowstone drill cores Y-1 (Honda and Muffler, 1970), Y-2 (Bargar and Beeson, 1981), and Y-13 (Keith and others, 1978a).

Figure 16 is a ternary diagram of acmite-jadite-hedenbergite + diopside (Ac–Jd–Hd+Di) electron microprobe compositions of aegirine from drill core Y-3 (table 5) compared with analyses of Y-2 hydrothermal aegirine (Bargar and Beeson, 1981). End-member compositions were calculated according to the method of Caithorn and Collerson (1974).

PECTOLITE

Pectolite is not an uncommon hydrothermal mineral, but, according to Browne (1978), this calcium-sodium silicate hydrate mineral is rare in active geothermal fields and has been identified only in the Y-3 drill core (at depths of 26.3-28.5 and 65.5-66.3 m) and in discharge pipes and channels of the New Zealand volcanic zone. In drill core Y-3, pectolite occurs as white fibrous crystals (figs. 15A, 15B, and 17A) or as radiating clusters of bladed crystals (fig. 17B) that were deposited later than aegirine and quartz.

FIGURE 14.—Scanning electron micrograph of a polycrystalline mat of lepidolite crystals from 28.3 m in drill core Y-3.
TRUSCOTTITE

Truscottite, another calcium silicate hydrate mineral, was identified in the Y-3 drill core from between 87.8 and 89.8 m, where the temperature measured during drilling was about 175 °C. (Truscottite has been formed in the laboratory from gyrolite and silica at 180 °C (Minato and Kato, 1967).) Clear, tabular truscottite crystals (fig. 18) were deposited in cavities and on fractures and are associated with earlier chalcedony, quartz, calcite, and laumontite, and later smectite and mordenite; Y-3 truscottite-bearing samples also contain chlorite and pyrite. Truscottite previously had been identified in three other Yellowstone drill cores (Y-2, Y-13, and C-I) (Keith and others, 1978a; Bargar and Beeson, 1981; Bargar and others, 1981). Several microprobe analyses of truscottite from 88.6 m in drill core Y-3 (table 6) suggest that this occurrence is slightly more calcium rich (chemical analyses of Y-3 drill-hole water compared with water from nearby Ojo Caliente Hot Spring, shown in table 1, suggest that calcium is precipitated during ascent of the water to the surface), is silica deficient, and has slightly greater alumina content than is found in most published analyses of the mineral (Minato and Kato, 1967; Juan and others, 1970) (fig. 19).

SULFATE MINERALS

GYPSUM

Gypsum occurs in a few scattered samples above 21.5 m in the Y-3 drill core. In the shallowest samples, white gypsum fibrous crystals line cavities and veins at less than 57 °C (the greatest temperature at which gypsum is the stable calcium sulfate phase) (Holland, 1967), and the mineral may be of hydrothermal origin. However, in deeper samples the gypsum is mostly deposited in association with pyrite and was probably formed by oxidation after the drill core was brought to the surface.
ALUNITE

Alunite, a potassium sulfate mineral, occurs in a single sample at 1.6 m in drill core Y-3 in association with opal, kaolinite, allophane, and halloysite. White and others (1975, table 3) indicate that the depth to the water table in drill hole Y-3 was about 2.4 m. Thus, alunite in Y-3 appears to have formed in the unsaturated zone of the drill core as a result of near-surface acid alteration, as described by Schoen and others (1974).

HEMATITE

Orange-red-reddish-brown iron oxide is sparse above 46 m and has a sporadic distribution throughout the remainder of the Y-3 drill core. Iron oxide (hematite where identified by X-ray diffraction) commonly occurs as pigment that stains chalcedony fracture fillings and vug deposits; however, occasional earthy or clayey hematite was noted. Hematite is an early hydrothermal mineral and appears to have formed later than chalcedony and pyrite, probably as a result of pyrite oxidation.

PYRITE

Tiny cubic pyrite crystals are found throughout much of the Y-3 drill core as deposits disseminated in groundmass or earlier green chalcedony. In Y-3, pyrite commonly lines fractures and vugs where the euhedral crystals or thin massive deposits are sometimes associated with later hematite. A few massive pyrite fracture fillings have crude hexagonal-appearing crystal cluster shapes that hint of a possible sulfide (pyrrhotite?) precursor. Pyrrhotite was identified in nearby Yellowstone drill core Y-2 (Bargar and Beeson, 1981). Y-3 pyrite crystals are typically small (about 0.1 mm); however, at about 4.5 m a thin, black, powdery pyrite deposit consists of 0.02–0.03 mm cubic crystals. At 45.5 m, pyrite

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**TABLE 5. -- Microprobe analyses of aegirine from Y-3 drill core**

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**Numbers of atoms on the basis of 6 oxygens**

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1 Sample numbers correspond to depth in meters.
SUMMARY AND DISCUSSION

The Y-3 drill hole penetrated about 42 m of surficial deposits consisting primarily of Pinedale glacial sediments that were deposited ≥45,000–14,000 years B.P. (Christiansen and Blank, 1974; Pierce and others, 1976). Below 42 m are two rhyolite flows, the Nez Perce Creek flow and an older, unnamed rhyolite flow, that Chris-
TABLE 6. -- Microprobe analyses of truscottite from 88.6 m in drill core Y-3

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<td>.11</td>
<td>.10</td>
<td>.05</td>
</tr>
</tbody>
</table>

¹Average of three analyses.

FIGURE 20.—Scanning electron micrograph of pyrite needlelike crystals deposited on top of quartz crystals that line a cavity at 45.5 m in drill core Y-3. Threadlike material may be mordenite. Smectite crystal clusters also appear to be present.

FIGURE 21.—Generalized paragenetic sequence of principal hydrothermal minerals in drill core Y-3. Length of lines is not indicative of duration of mineral deposition.
tiansen and Blank (1974)correlated with the Central Plateau Member of the Pleistocene Plateau Rhyolite. The Nez Perce Creek flow has K-Ar dates between 140,000 and 160,000 years B.P., as reported in White and others (1975), Waldrop and Pierce (1975), and Leeman and Phelps (1981). Other rhyolite flows of the Central Plateau Member were extruded between about 200,000 and 70,000 years ago (Christiansen and Blank, 1972). Thus the unnamed rhyolite flow in drill core Y-3 is probably not much older than the overlying Nez Perce Creek flow.

The effects of hydrothermal alteration are evident throughout the Y-3 drill core. Silica and zeolite minerals provide the cementing material for much of the surficial deposits. Parts of the two rhyolite flows were bleached by thermal waters, and the entire drill core contains hydrothermal minerals lining vugs and filling fractures. A generalized paragenetic sequence for the principal hydrothermal minerals found in drill core Y-3 is shown in figure 21. The glacial sediments and both rhyolite flows probably have been subjected to hydrothermal alteration throughout their histories, because the duration of hydrothermal activity in the Yellowstone region appears to extend over the last 266,000 years and may possibly date back to about 600,000 years B.P. (Silberman and others, 1979). Consequently, some of the hydrothermal minerals in drill core Y-3 probably predate the Pinedale glacial sedimentary deposits of the upper 42.2 m. Chalcedony, pyrite, hematite, quartz, and clay minerals (smectite and mixed-layer illite-smectite) are early hydrothermal deposits in the volcanic section of the drill core (listed in order of deposition); calcite, chlorite, fluorite and zeolite minerals are later deposits, although the depositional sequence of calcite, chlorite, and fluorite was not determined. Chalcedony, pyrite, hematite, quartz, and clays are also shown as late hydrothermal deposits in figure 21 because these minerals are found in the younger sedimentary section of the drill core, as well as in the volcanic section. In addition, pyrite was deposited both earlier and later than quartz, and quartz occurs both earlier and later than calcite in the same sample.

Porosity, based on the difference between bulk- and powder-density measurements, was computed for 16 selected samples of Y-3 drill core. The calculated porosity ranged from 15.7 to 45.8 percent (average 35.1 percent) in the glacial sediments and from 3.7 to 36.9 percent (average 15.3 percent) in the rhyolite flows. The porosity and effective permeability of the volcanic rocks appear to have been greatly reduced owing to precipitation of hydrothermal minerals, particularly quartz and chalcedony, in vugs and fractures. In fact, many of the fractures in the lower, unnamed rhyolite flow have been completely closed (self-sealed) by silica deposition.

According to Pierce (1979, pl. 1), as much as 600 m of Pinedale glacial ice may have covered the ground surface about 10 km north of the Y-3 drill hole. In the thermal areas of Lower Geyser Basin, a significant portion of the ice probably melted and formed an ice-dammed lake (Muffler and others, 1971), the possible extent of which is shown by Pierce (1979, fig. 35). Muffler and others (1971) suggest that the triggering mechanism for the nearby hydrothermal explosion crater at Pocket Basin (fig. 1) may have been a sudden release of pressure caused by breaching of the ice dam and rapid draining of the lake. Prior to failure of the dam, the confining pressure at a given depth beneath the ground surface would have greatly exceeded present-day hydrostatic or lithostatic pressures, and the temperature at that depth would have been significantly warmer than the temperature measured during drilling of Y-3. Presumably, the fluid-inclusion homogenization temperature measurements for the Y-3 drill core (and other drill cores in Lower Geyser Basin) that exceed the temperatures measured during drilling reflect the higher temperature-pressure conditions that existed as a result of increased overburden during the Pinedale glacial time (Bargar and others, 1984).

In the sedimentary section of the Y-3 drill core, hydrothermal minerals containing sodium as a dominant cation (analcime, clinoptilolite, mordenite, Na-smectite, and aegirine) are more abundant than calcium-bearing minerals (calcite, fluorite, Ca-smectite, and pectolite). Conversely, in the volcanic section of the drill core, calcium-rich minerals (dachiardite, laumontite, yugawaralite, calcite, fluorite, Ca-smectite, pectolite, and truscottite) are predominant over sodium-bearing minerals (aegirine, mordenite, and Na-smectite). A few hydrothermal minerals that contain significant amounts of potassium (alunite, illite-smectite, and lepidolite) occur in both sections of the drill core.

Rhyolite flows of the Central Plateau Member are similar in chemical composition to the Summit Lake flow (Keith and others, 1978b) and to glass from the Nez Perce Creek flow (Leeman and Phelps, 1981). The analyses of the latter two flows show abundant sodium and potassium and little calcium. Twelve hydrothermally altered rocks from the Nez Perce Creek flow and the underlying unnamed rhyolite flow were analyzed by rapid rock methods and reported by Beeson and Bargar (1984). Figure 22 shows A-F-M and Na₂O-K₂O-CaO diagrams for these Y-3 volcanic rocks, as well as for hydrothermally altered glacial sediments from the upper part of the Y-3 drill core. All of the rocks are alkali rich (especially potassium) and calcium poor. As such, the dominance of calcium-bearing hydrothermal minerals in the lower rhyolitic volcanic section of Y-3 would appear to be due to the waters that deposited the miner-
als. In fact, a comparison of chemical analyses of water from the Y-3 drill hole and from nearby Ojo Caliente Hot Spring (table 1) suggests that calcium, along with potassium, is probably lost during adiabatic cooling of an ascending boiling water, whereas sodium remains unchanged after correcting for boiling (Bargar and others, 1973).

The silica geothermometer for Ojo Caliente Hot Spring water (176°C) is in good agreement with the measured temperature (174°C) for water from the 88-m aquifer in drill hole Y-3 (table 1, column 2). Other geothermometer calculations in table 1 give temperatures that are too low for Y-3. Calculated geothermometers for the bottom-water (table 1, column 1) are significantly lower than the 196°C measurement at the bottom of the drill hole. At the time the Y-3 bottom water sample was collected, White and others (1975) concluded that the region below 80 m in the drill hole was influenced by downflow of cooler water from the upper part of the drill hole.

Fluid-inclusion homogenization temperatures were measured in quartz (48 fluid inclusions in 4 samples) and fluorite (47 fluid inclusions in 3 samples) crystals from the Y-3 drill core. Most of the liquid-rich fluid inclusions appear to be secondary(?) or pseudosecondary and have homogenization temperatures that plot very close to the present-day measured temperature curve (fig. 23). One homogenization temperature measurement of 313.5°C at 45.8 m was not reported in figure 23 because the value obviously was too high owing to necking-down of the fluid inclusion by precipitation of later silica (Roedder, 1984). Similarly a few other small, irregular fluid inclusions, some with “tails,” may give erroneous (either too high or too low) homogenization temperatures. Probably the most reliable homogenization temperatures shown in figure 23 are the clustered measurements, which usually plot less than about 20–40°C from the measured temperature curve. From these data we infer that the temperature of the water that deposited the quartz and fluorite crystals studied was a little warmer than the temperatures measured during drilling.

REFERENCES CITED

