

**Mahd adh Dhahab: Precambrian
epithermal gold deposit in Saudi Arabia**

by

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U.S. Geological Survey

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ABSTRACT

The largest and most productive mine in Saudi Arabia in ancient and modern times has been Mahd adh Dhahab. Country rocks in the district are pyroclastic and clastic rocks which belong to the Halaban Group of late Precambrian age. From oldest to youngest, the units are: andesite, lower agglomerate, lower tuff, upper agglomerate, and upper tuff. These units have been tilted northerly forming a homocline and are complexly broken by six sets of faults; three of these sets, which strike $N 25^{\circ}-30^{\circ} W$, $N 10^{\circ} W$ to $N 20^{\circ} E$, and $N 30^{\circ}-60^{\circ} E$, contain productive quartz veins.

The principal ore minerals are native gold and silver; pyrite, chalcopyrite, sphalerite and galena, plus minor tetrahedrite and argentite. The gold and silver is finely disseminated in the veins and in the altered selvages of the veins. Widespread potassic and propylitic alteration accompanied ore formation.

Early widespread propylitic and potassic alteration and sulfide and quartz precipitation are believed to have taken place at temperatures below $350^{\circ}C$, based on K feldspar stability and S isotope data. Most of the quartz and precious metals precipitated in the range $200^{\circ}-150^{\circ}C$, based on studies of fluid inclusions. The isotopic data indicate that sulfur and carbon came from a deep seated source, but that the water, especially in the later stages, was meteoric.

INTRODUCTION

1 Mahd adh Dhahab, literally "cradle of gold", mine was the most productive
2 mine in Saudi Arabia in both modern and ancient times. It is located roughly
3 midway between Makkah and Medina, 275 km northeast of Jiddah on the western
4 edge of the Arabian Shield (Fig. 1). Modern investigation of the mine began
5

6 Figure 1 near here.

7
8 in 1935 by K. S. Twitchell, an American mining engineer, whose evaluation
9 led to the formation of the Saudi Arabian Mining Syndicate (SAMS). Production
10 by SAMS during the years 1939 to 1954 was large, totaling 765,768 fine ounces
11 of gold and 1,002,029 ounces of silver (Goldsmith and Kouther, 1971). A quarry
12 level, and 4 underground levels to a depth of 200 meters (m), were worked.
13

14 It was estimated by Dolph (1941) that nearly 1,000,000 metric tons of
15 tailings were piled in dumps around the ancient workings. Ancient production
16 must have been great in this largest ancient mine of Saudi Arabia. Tens of
17 thousands of stone implements used for ore dressing and mining still litter
18 parts of the mining area. In his book on Saudi Arabia, Twitchell (1958) noted
19 two distinct layers of ancient tailings; the younger tailings were dated by Kufic
20 inscriptions as belonging to the Abbasid Caliphate (750-1258 A.D.).
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Figure 1.--Index map showing the location of the Mahd adh Dhahab district.

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 (T) should be added here
 about the 1:5000
 region with Saudi Arabia. The
 boundary
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 1:5000

The geology of Jabal Mahd adh Dhahab and the immediate vicinity was mapped at a scale of 1:2400 in 1972-73 (Luce, Bagdady, and Roberts, 1976).

Fig. 2 is a simplified version of that map of the 3-km² area which includes

Figure 2 near here.

the SAMS quarry in the northeast corner. Data mapped on plane table and aerial photographic bases were compiled with a stereoplotter and incorporated a 1:480 scale map made by Dirom (1947) of the SAMS mine area in the northeast corner. Coordinated with the geologic investigation was an extensive surface geochemical sampling program (Roberts, Bagdady and Luce, 1978). A subsequent drilling program and more detailed surface mapping in the eastern third of the area was done under the direction of Ronald Worl of the U.S. Geological Survey. Partly as a result of interest generated by these studies, Consolidated Gold Fields, Ltd. obtained a license to the district in 1975.

REGIONAL GEOLOGY

Jabal Mahd adh Dhahab rises to an altitude of 1238 m above the 1000 meter elevation of the wadis which drain to the north and south of the area shown in Fig. 2. The jabal (mountain) protrudes above the great Najd pediplain which slopes gently northeastward from the Red Sea Escarpment.

The general regional geology of the northwest corner of the 4⁰ Southern Hijaz quadrangle, wherein Mahd adh Dhahab lies, is that of northeast trending Precambrian greenstones, volcanic and clastic sedimentary rocks, complexly folded Precambrian schists and Precambrian granitic intrusions of several ages (Brown et al, 1960).

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Figure 2.--Geologic map of Mahd adh Dhahab district, Kingdom of Saudi Arabia.

1 The same general types of rocks are found in the immediate region surrounding
2 Mahd adh Dhahab (Goldsmith, 1971; Goldsmith and Kouther, 1971). Precambrian
3 volcanic rocks of rhyolitic and andesitic composition, which are dominantly
4 clastic in origin, are moderately folded along northeast axes. A large northeast
5 trending right-lateral fault system passes east of Mahd adh Dhahab. Subsidiary
6 block faulting has affected Jabal Mahd adh Dhahab. Basaltic dike swarms
7 which trend north to northwest south of the mine area have apparently been
8 shifted to a northeast trend in the vicinity of the fault system.

9 SEDIMENTARY AND VOLCANIC ROCKS

10 Taken as a whole, the predominantly pyroclastic rocks in the Mahd adh
11 Dhahab area appear to have been deposited on the flanks of a caldera, although
12 no direct evidence of such a structure remains today. By lithologic correlation
13 with other areas, we believe them to belong to the Halaban Group (Greenwood
14 et al, 1976).

15 The sequence of rocks of late Precambrian age in the Mahd adh Dhahab
16 area from oldest to youngest is andesite, lower agglomerate, lower tuff, upper
17 agglomerate, and upper tuff. During a study of the regional geology in 1966,
18 Goldsmith and Kouther used Maclean's (1961) terminology, Mahd adh Dhahab
19 series, and correlated the series with the Murdama and Fatima Formation.
20 More recently, Aguttes and Duhamel (1971) have assigned the Mahd adh Dhahab
21 series to the next older Halaban Group.

Andesite

The oldest rock unit in the area is andesite which crops out in an east-west belt in the southern part of Jabal Mahd adh Dhahab and on pediments south of the jabal. Goldsmith and Kouther (1971) suggested that the andesite is unconformably underlain by the younger units in the jabal, but this relationship cannot be seen in the map area due to faulting along the boundary. The thickness of the andesite was not measured, but is estimated to be more than 300 m.

The andesite is generally fine grained, although coarser porphyritic layers are locally present. Some units are crystalline and resemble diabase or diorite. A few layers are fragmental, suggesting a pyroclastic origin. The combination of these diverse lithologic units suggests that the andesite is partly volcanic, partly intrusive, and partly pyroclastic in origin.

Microscopically, the andesite shows a pilotaxitic texture of fine plagioclase laths in a matrix of chlorite and clay minerals. The rock is so propylitized that the composition of the plagioclase could not be determined and no original mafic minerals remain because of strong alteration to chlorite, epidote, and magnetite.

Lower agglomerate

The lower agglomerate unit crops out on the south flank of Jabal Mahd adh Dhahab. The contact with the underlying andesite is a normal fault that cuts east-west diagonally across the agglomerate, causing a gradual narrowing of the belt of outcrop.

1 The lower agglomerate is a lapillistone and pyroclastic
2 breccia with a tuffaceous matrix. Lithic fragments
3 in basal units are mostly subangular andesite; fragments of
4 rhyolitic composition are locally abundant in the upper
5 part. Bedding and sorting is poor for the most part
6 and attitudes can be taken only on thin tuffaceous
7 interbeds. The lower agglomerate, like the upper agglomerate,
8 is a lahar deposit. Epidote and chlorite are common
9 alteration products of the matrix.

10 The lower agglomerate is extensively mineralized
11 along faults in the southeastern and south-central
12 parts of the jabal. The agglomerate was apparently
13 competent and maintained open fractures during metallization,
14 permitting the formation of mineralized quartz veins
15 and silicification of adjacent rock.

16 Lower tuff

17 The lower agglomerate grades upward into the lower
18 tuff through an interbedded zone about 10 m thick.
19 The lower tuff is a cliff-forming, resistant unit that
20 makes up the backbone of Jabal Mahd adh Dhahab and
21 forms the highest point.
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1 The lower tuff is mostly a fine-grained and laminated, indurated siliceous
2 ashfall tuff with some sandy units. A stratigraphic section was measured in
3 the central part of the jabal where the unit is about 207 m thick. This section
4 may be conveniently divided into three parts: a lower laminated ashfall tuff;
5 a middle sandstone and graywacke; and an upper interlayered tuff, shale, and
6 lapilli crystal tuff.

7 The lower part, 55 m thick, is characterized by thin-bedded, fine-grained,
8 felsic ashfall tuff which is interbedded with lithic lapillistone. The tuff contains
9 andesitic fragments, sanidine shards and devitrified spherulites, now partly
10 chloritized, and potassium feldspar crystal tuff layers. Near the top of this
11 lower part, the lapilli tuff component becomes less prominent and the tuff
12 becomes thicker bedded.

13 The middle part of the lower tuff unit, 72 m thick, is characteristically
14 medium- to fine-grained sandstone and subgraywacke with lapillistone layers
15 in places. Beds near the upper boundary of this part are commonly stained
16 with iron oxides which form prominent liesegang rings.

17 The top part, 80 m thick, is made up of alternating tuff, sandstone, rare
18 shale beds, and local lapilli crystal tuffs. The uppermost 28 m is strikingly
19 hornfelsic.
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1 The lower tuff locally contains small veins along faults, but in general
2 the lower tuff was not extensively mineralized. The largest mineralized areas
3 in this unit are in the southeastern and western parts of the district.

4 Upper agglomerate

5 The upper agglomerate rests conformably on the lower tuff. Measure-
6 ments on cross sections give an average thickness of about 140 m.

7 The upper agglomerate unit is similar in general appearance to the lower
8 agglomerate; it is a crystal lithic lapillistone with subangular tuff, jasper,
9 and andesitic lithic fragments.

10 The upper agglomerate, like the lower agglomerate, was competent,
11 fractured well, and maintained open fractures. These characteristics made
12 it a favorable host for metallization and the fractures are now filled with
13 quartz.

14 Upper tuff

15 The youngest stratigraphic unit in the district is the upper tuff which
16 crops out in a narrow east-west zone on the northern edge of the jabal; it dips
17 north 45° - 55° and underlies the wadi, where it is largely covered by a thin
18 layer of alluvium and mine dumps.

1 The upper tuff is a thin-bedded, fissile, ashfall tuff. In places, it contains
2 lapillistone layers 10 to 20 cm thick. Microscopically, it is seen to consist
3 of ashfall tuff with some introduced potassium- and plagioclase feldspars,
4 quartz, and calcite.

5 The upper tuff rests unconformably on the upper agglomerate unit. Many
6 quartz veins appear to terminate against the upper tuff, but this may be because
7 the tuff was a relatively impermeable barrier to hydrothermal solution movement.

8 IGNEOUS ROCKS

9 Andesite dikes

10 A series of andesite dikes up to 2 m wide were emplaced along north-
11 easterly-trending fault zones in many parts of the jabal.

12 The dikes are composed mostly of saussuritic aggregates of altered
13 plagioclase, and chlorite which was largely derived from plagioclase and mafic
14 minerals.

15 Some dikes contain amygdules, mostly composed of chlorite, calcite,
16 and quartz. The dikes were emplaced late in the ore-forming process as they
17 cut and displace veins in many areas. Metallization continued on a small scale
18 after dike emplacement as some gold- and silver-bearing quartz-calcite veins
19 cut the dikes locally. Analyses show that some of these late veins contain
20 small amounts of gold and silver.

SURFICIAL DEPOSITS

Three surficial units--talus, older alluvium, and younger alluvium--were mapped (Luce, Bagdady, and Roberts (1975). These are omitted from Fig. 2 for clarity. Talus consists of coarse debris that mantles the lower slopes of the jabal. It includes material of a wide age span from Late Tertiary or Early Quaternary to Holocene and also includes dumps discarded by ancient and modern miners.

Older alluvium includes fan or terrace gravels which are commonly covered by a coating of desert varnish. The finer fractions generally have been removed by wind and water, leaving a lag pebble pavement.

The younger alluvium is found in the present wadis. It is commonly lighter in color than the older alluvium because it has been reworked during the torrential floods that occur several times a year in this area. Some areas underlain by younger alluvium contain placer gold deposits, but the richer ores are commonly in the lower gravel layers, which probably are part of the older alluvium. Placer and eluvial gold deposits were mined by ancient miners on the flanks of Jabal Mahd adh Dhahab and in valleys that drain the jabal.

STRUCTURE

The principal structural feature at Madh adh Dhahab is an east-trending homocline that dips northerly (fig. 2). The homocline in the southern part of Jabal Mahd Dhahab dips northerly 30° - 40° . The dip gradually steepens in the central-part to 50° - 60° N and in the northern part to 70° - 75° N. The homocline is complexly broken by six sets of faults which are listed below, numbered in order of formation:

<u>Fault set</u>	<u>Trend</u>	<u>Mineralization</u>
6. N 45° W; vertical to steeply SW		None
5. N 30° - 45° E; steep to vertical		Contains andesite dikes
4. N 30° - 60° W; steeply SW to vertical		Locally significant
3. N 10° W to N 20° E; steeply W		Major
2. N 30° - 60° E; 35° - 65° NW, steepening		Major
1. N 25° - 30° W; 55° - 65° NE to 70° SW		Minor

1 The oldest fault set (1) trends $N25^{\circ}-30^{\circ}W.$, has been
2 broken and displaced by other sets, and is so fragmented
3 that only short segments are now seen. Ancient mine
4 workings along faults of this set indicate small-scale
5 mineral production.

6 The second fault set strikes $N30^{\circ}-60^{\circ}E$ and commonly
7 dips $35^{\circ}-65^{\circ}NW$; many mineralized veins and silicified
8 zones are along faults of this set. Quartz veins on
9 this set tend to be lenticular and show shearing along
10 their margins as well as internally. Their generally
11 low dips suggest that they may be thrust faults. The
12 west silicified zone forms a series of sheared flatirons
13 up to 4 m thick that cap large areas of the slope on
14 the western side of the jabal. These flatirons have
15 been cut by north-south faults of the third set and cut
16 and displaced by the fourth set of faults which strike
17 $N30^{\circ}-60^{\circ}W$ and dip steeply.
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1 The third fault set strikes north-south (N20°E
2 to N10°W) and dips steeply west. Normal faults of
3 this set offset the contact of the lower agglomerate
4 and lower tuff in a series of steps; the downthrown
5 block is generally on the west. This set also contains
6 major mineralized quartz veins which strike north to
7 N10°E. Veins in this set of faults have been extensively
8 mineralized, especially in the lower and upper agglomerates
9 at the northeastern end of the jabal. Two veins of
10 this set have been productive in underground workings.
11 There has been repeated movement along faults of this
12 set, and quartz fillings are commonly brecciated.

13 Long, continuous faults which strike N30°-45°E
14 form the fifth set. Many of these show considerable
15 left-lateral displacement. Two faults at the eastern
16 edge of the jabal strike north-northeast but have right-
17 lateral displacement and may be part of this set.
18 Andesite dikes in this northeast fault zone attest
19 to later reactivation. These are not zones of major
20 quartz veining; however, young brecciated quartz veins
21 follow the attitude of the dikes which they cut.

1 The sixth and youngest fault set strikes northwest
2 and dips vertically to steeply southwest. These are
3 small transverse faults which cut the third fault-vein
4 set and generally show left-lateral offsets of as much
5 as a meter or two. They are probably related to regional
6 Najd faults, which are discussed later.

7 The faults at Mahd adh Dhahab formed in response
8 to regional stresses during several periods. Shearing
9 along northwest and northeast directions, with a probable
10 north-south principal stress, is inferred during the
11 formation of faults of the first and second sets.
12 A continuation of the same general stress field is
13 thought to have produced the fourth set. The north-
14 south faults of the third set are uncommon in the region
15 outside Jabal Mahd adh Dhahab and most likely originate
16 from vertical uplift. Faults of the fifth set show
17 progressive downthrow on the west and may be related
18 to uplift of the jabal during and following regional
19 emplacement of intrusive bodies as well as a continuation
20 of the general stress field. Faults of the sixth set
21 seem to be related to northwesterly movements during
22 Najd wrench faulting late in Precambrian time; one
23 strand of the Najd fault system passes a few kilometers
24 north of the district.

METASOMATISM AND QUARTZ VEINS

Intense silicic and potassic metasomatism is evident in the area of SAMS mining on the northeastern edge of the map area in the upper agglomerate and also on the western tip of the jabal in the lower agglomerate. Both of these units in the two areas are generally silicified and the rock matrix recrystallized. Primary quartz and K-feldspar phenocrysts have been embayed and generally fractured. The K-feldspars have been remade to a more potassium-rich composition not only in these two large areas but also in many coarse and therefore permeable layers within the relatively lower tuff and in both agglomerate units. There has also been an introduction of K-feldspar in early bands of some quartz veins, especially in the SAMS mining area. Sericitic alteration is found in the same places as the potassic alteration but occurs to a minor degree. Argillic alteration is slight and it is not yet known to what extent feldspar alteration to clays is caused by weathering.

Two feldspar separates, one of a banded ore vein and one of metasomatized phenocrysts from a thin sill in the lower tuff, have been analyzed by the x-ray diffraction method of Wright (1968) and Wright and Stewart (1968). They are nearly identical maximum microclines with a composition of Or_{100} . The upper stability limit of maximum microcline at $P_{\text{H}_2\text{O}}$ less than 0.5 kb, a reasonable estimate for Mahd adh Dhahab, is 375° (Tomisaka, 1962; Wright, 1967) so it is likely that the vein feldspars and the chemically altered phenocrysts formed below this temperature. The age of the vein feldspar has been determined by Zell Peterman (written commun. from R.J. Fleck, 1973). Assuming an initial $^{87}\text{Sr}/^{86}\text{Sr}$ ratio of 0.704, the data yield an age of about 675 m.y., making the hydrothermal alteration and metallization late Precambrian in rocks that correlate with late Precambrian rocks.

The intensely metasomatized areas were originally mapped as rhyolite porphyry (Luce et al, 1975) but upon further examination and discussion (Worl, personal commun., 1977) it is believed that most rock textures are pyroclastic. There are units within these areas that show flow banding and sanidine phenocrysts, but many of these can be shown to be lithic fragments. However, certain features such as doming of strata underground in the SAMS mine area (Dirom, 1947) and recrystallization may be related to a near-surface intrusive, as yet not found.

1 The great preponderance of introduced material
2 is in quartz veins, with which metallization is associated.

3 Four types of quartz veins are recognized at Mahd
4 adh Dhahab on the basis of mineralogy, quartz crystallization,
5 and structure. The first three types are older than
6 the andesite dikes, and the fourth type is younger
7 than the dikes.

8 Type 1 veins are mainly massive, milky white quartz
9 which formed early in the metallogenic cycle. These
10 veins contain sparse pyrite and commonly have a low
11 gold content.

12 Type 2 veins are composed of banded and crustified,
13 milky to clear quartz, generally with comb or cockade
14 structure. Red to pink potassium feldspar (maximum
15 microcline) is commonly an early mineral in this type
16 of vein, and was followed by banded base metal sulfides-
17 chalcopryrite, galena, and sphalerite. Fe-rich chlorite
18 is commonly interlayered with the sulfides and quartz.

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2 Chemical analyses were made of some type 2 veins
3 to determine the distribution of gold and silver in
4 them. Figure 3 shows one of these veins. Silver is

5
6 **Figure 3 near here.**

7
8 distributed throughout the specimens in galena and
9 tetrahedrite, but gold is mainly in the selvage zones.
10 The two generations of quartz in the specimen shown
11 contain comparatively little gold. According to Goldsmith
12 (1971) much of the gold mined by SAMS was in the selvages
13 of quartz veins; microscopic study of ore specimens
14 supports his statement.
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Figure 3.--Type 2 quartz vein from lower agglomerate showing
crustification structure. Chemical analyses are of the different
banded units.

1 Type 2 veins show well developed banding, cockade,
2 and crustified textures which testify to formation of
3 the vein at shallow depths in open fractures (Fig. 3).
4 In some type 2 veins, base metal sulfides are especially
5 abundant and may constitute a base metal ore.

6 Type 3 veins are characterized by comb quartz,
7 sphalerite and chalcopryrite, and a late stage carbonate
8 such as ankerite, manganiferous calcite, or calcite.
9 Banded veins showing repeated reopening and filling
10 textures are common (Fig. 4).

11
12 Figure 4 near here.

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14 Quartz veins of type 4 which cut the andesite
15 dikes are commonly calcareous and contain only sparse
16 sulfides, mostly pyrite plus minor chalcopryrite.

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Figure 4.--Disseminated sphalerite (Sl) + galena (gn)
in chloritic gangue bordered by quartz veins,
chlorite layers, and late-stage massive sphalerite
(Sl₂).

Ore minerals

The primary sulfide minerals in order of abundance are pyrite, chalcopyrite, sphalerite, galena, tetrahedrite, and argentite. Native gold and silver accompany the major sulfides. The sulfide minerals were mostly crystallized in open spaces and are commonly euhedral. The early pyrite is generally in the form of cubes as much as 0.5 cm on a side. Chalcopyrite commonly replaces pyrite along fracture zones and is mostly fine grained, although some late-stage chalcopyrite is coarsely crystalline. Sphalerite ranges from finely to coarsely crystalline and dark gray to black. It commonly contains exsolved chalcopyrite in a emulsion texture. Galena is generally fine grained and is closely associated with chalcopyrite and sphalerite. Tetrahedrite and chalcopyrite are frequently found altered in part to covellite and malachite. Native silver is very fine grained and is mainly associated with galena, sphalerite, and tetrahedrite. Minor argentite and a mineral suspected to be pyrargyrite or polybasite have also been found. Gold is most commonly fine grained and occurs as the free metal in quartz or chlorite. A few small crystals and some gold wires were seen in hand specimens, but generally the grains are too small to be recognized except under the microscope. The native gold sometimes has tiny inclusions of a weakly anisotropic mineral which is probably a telluride such as calaverite.

1 The sulfide minerals are commonly weathered in the oxidized zone to
2 a depth of 10 meters or more. Pyrite alters to limonite and hematite; chalcopyrite
3 and tetrahedrite to covellite, chalcocite, malachite, and chrysocolla; sphalerite
4 to hydrozincite, hemimorphite, and smithsonite; galena to anglesite and cerussite
5 the mangiferous carbonates to wad and pyrolusite.

6 PARAGENESIS

7 Study of polished slabs, ore mounts, and specimens collected from open
8 stopes and dumps has established the general paragenesis of ore and gangue
9 minerals; the sequence of these minerals is shown in Table 4.

10 NATURE OF THE MINERALIZING SOLUTIONS

11 Standard techniques for fluid inclusion and isotopic study were employed
12 on specimens from drill core, outcrop, and from the SAMS mine dump mainly
13 taken from the highly mineralized eastern third of the jabal. Variations in
14 the isotopic compositions and fluid inclusion temperatures are slight within
15 the small area and depths (≤ 200 m) sampled indicating that the conditions
16 of hydrothermal alteration and mineralization were fairly uniform with an
17 expected general but slight increase in temperature with depth.
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Fluid inclusions

The vein quartz contains a great abundance of tiny secondary fluid inclusions as a result of entrapment of liquid following the intense fracturing in the district. More scarce large ($< 20 \mu\text{m}$) primary and pseudosecondary fluid inclusions in diverse veins gave homogenization temperatures which spanned the range 142° to 278°C . Salinity was quite low in all samples measured, and ranged from 0 to 0.1% equivalent NaCl by weight. Several of the higher temperature inclusions (254° - 278°C) contained liquid CO_2 with a maximum volume proportion estimated at 40%.

Sulfur isotopes

The $\delta^{34}\text{S}$ values of sulfides are typical of post-magmatic vein-type hydrothermal deposits (Table 2). For individual minerals, values vary within a narrow range: from 2.7 to 3.2‰ in pyrite and from 0.8 to 2.6‰ in sphalerite, but probably cover a wider range in the paragenesis. $\delta^{34}\text{S}$ values in coexisting mineral pairs (gn-sl, py-sl, and py-cpy) indicate temperatures of 210°-290°C. All of these minerals probably precipitated prior to the deposition of late vein quartz and thus the indicated temperatures are reasonable for depositional temperatures. The fact that the mineral pairs always give reasonable temperatures even though they are not contemporaneous suggests that the chemistry of the hydrothermal solutions was very tightly controlled during the deposition of the various sulfides by equilibration with the volcanic host rocks in the ore zone or at depth. It is this tendency toward equilibration, along with the lower $\delta^{34}\text{S}$ values, that separates the Mahd adh Dhahab sulfur isotope data from that for Jabal Sayid (50 km northwest) and other massive sulfides in Arabia (Rye, Roberts and Mawad, in prep.).

The sulfur at Mahd adh Dhahab may have been derived from igneous sources, or possibly from a massive sulfide deposit, when one considers the average $\delta^{34}\text{S}$ value of many of the Arabian massive sulfides.

Oxygen, carbon, and hydrogen isotopes.--Field and petrographic observations that carbonate formed later than most of the quartz are substantiated by the ^{18}O disequilibrium between coexisting quartz and carbonate (Table 3). Calcite concentrates ^{18}O relative to quartz in all cases except sample 97896 for which $\Delta_{\text{Q-C}} = +1.7\%$.^{1/} This case corresponds to a temperature

^{1/} Subscripts are Q, quartz; C, carbonate; W, water; Chl, chlorite; M, muscovite. $\Delta_{\text{Q-C}} = \delta^{18}\text{O}_{\text{quartz}} - \delta^{18}\text{O}_{\text{calcite}}$

of 450°C (Bottinga and Javoy, 1973) which is not in agreement with all other temperature determinations. A more reasonable explanation is that carbonate formed later and at a lower temperature than quartz. This is the obvious explanation where $\delta^{18}\text{O}$ is +15 or +19‰.

The $\delta^{18}\text{O}$ values of the water involved with quartz ranges from -0.3 to 3.1‰, if we take $T = 200^\circ\text{C}$ for the formation of all quartz analyzed ($\Delta_{\text{Q-W}}^{200} = 12.0\%$, Clayton et al., 1972). The exception is 64105 quartz with a $\delta^{18}\text{O}$ value of 15.66‰. If 64105 quartz formed at 200°C, $\delta^{18}\text{O}_\text{W} = +3.56\%$, and might indicate a magmatic component.

More likely it could have formed in water of $\delta^{18}\text{O} \approx 0\text{‰}$, but at a temperature $\approx 150^\circ\text{C}$.

Likewise, both microcline types, which have similar $\delta^{18}\text{O}$ values, thus suggesting a similar origin, yield a range of $\delta^{18}\text{O}_\text{W} = -0.1$ to 1.0‰ using the equation of Bottinga and Javoy (1973). A similar value for the water causing the chloritization can be obtained by a circuitous route. Primary Sierra Nevadan chlorites have an average $\delta^{18}\text{O}$ value of $+4.0\text{‰}$ (O'Neil and Dodge, unpub. data) and with $\delta^{18}\text{O}_\text{W}$ assumed to be $+8.0\text{‰}$, a typical magmatic value, we reason to a $\Delta_{\text{Chl-W}}$ value of -4.0‰ at 700°C . Now, $\Delta_{\text{M-W}} = -1.0\text{‰}$ at the same temperature and $\Delta_{\text{M-W}} = 3.0\text{‰}$ at 200°C (Bottinga and Javoy, 1973). If we assume the change in the isotopic fractionation factor with temperature, $\frac{\partial \alpha}{\partial T}$, to be approximately the same for chlorite-water and muscovite-water, we conclude that $\Delta_{\text{Chl-W}} = +3.0\text{‰}$ at 200°C . Therefore, water in equilibrium with 69026 chlorite would have had $\delta^{18}\text{O} \approx -0.4\text{‰}$. The assumptions involved in this calculation are not so gross as to affect the value more than $1\text{--}2\text{‰}$.

Thus, all indications are that the hydrothermal waters had $\delta^{18}\text{O}$ values near zero and probably up to 3‰ lighter. Despite the fact that the Mahd adh Dhahab hydrothermal solutions were Precambrian, they were isotopically similar to the only modern ground waters measured in the area (Oman) for which $\delta^{18}\text{O}$ ranges from +0.4 to -1.7‰ (Barnes et al, 1978). This similarity, coupled with the low salinity in fluid inclusions, leads us to the conclusion that the waters were preponderantly meteoric. The mineralizing waters of Mahd adh Dhahab closely resemble, in isotopic composition and low salinity, those of epithermal Tertiary Au-Ag deposits in the Great Basin of Nevada (O'Neil and Silberman, 1974).

The carbon isotope data is all in the narrow range of -7.58 to -5.66‰, which is classical for mantle-derived carbon (Taylor et al, 1967; Pineau et al, 1976). Again this resembles younger epithermal deposits in the western U.S. where carbon and sulfur had a deep-seated source and water at least in the late stages, had come from above.

Deuterium isotope data are more ambiguous. Concerning the chlorites $\Delta_{\text{Chl-W}}$ at $200^{\circ}\text{C} \approx -70\%$, based on minerals such as serpentine and muscovite. The exact fractionation depends on the chemical composition of the octahedral site (Suzuoki and Epstein, 1976). Therefore, it appears that if $\delta D_{\text{Chl}} = -65\%$, then the water had a δD_{W} value close to zero, compatible with the oxygen isotope calculations. Quartz samples 64105 and 64120 gave δD values of -75 and -39% , respectively, for the inclusion waters. Chlorite could have formed in water of $\delta D = -39\%$ if the temperature were about 400°C . δD_{Q} for sample 64105 remains anomalous for being so light, but perhaps there was a magmatic component to its fluid.

Summary

The Precambrian pyroclastic rocks that make up Jabal Mahd adh Dhahab range from andesitic to rhyolitic composition and were probably deposited in a caldera environment. They were tilted but only gently folded into a homoclinal structure which was then repeatedly sheared, fractured, and block faulted. Silicic and potassic metasomatism took place in the northeastern and western parts of the jabal and also formed northeast, northwest, and north-trending quartz veins throughout. An epithermal mineral assemblage consisting mainly of pyrite, chalcopyrite, sphalerite, galena, tetrahedrite, argentite, tellurides, and native gold and silver was introduced into the quartz veins and their chloritic selvages. Age dating of vein feldspar shows that mineralization took place in late Precambrian time.

1 The early hydrothermal solutions which propylitically
2 altered the country rock and precipitated K-feldspar,
3 sulfides and some quartz had temperatures in the range
4 200°-310°C. Later solutions which precipitated quartz
5 and precious metals had temperatures in the range 150°-
6 200°C and had low salinities. Introduced sulfur and
7 carbon came from a deep seated source while most of
8 the water, particularly in the later stages had a meteoric
9 origin. The Precambrian hydrothermal solutions have
10 a similar isotopic composition to present day ground
11 water in the region, an interesting but perhaps fortuitous
12 fact.

13 The mechanism for preservation of this high crustal
14 level mineralization in the Arabian Shield probably
15 had its origin in block faulting of the jabal.

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Table 1. Paragenetic sequence of primary ore and gangue minerals, Mahd adh Dhahab.

	Type 1 veins	Type 2, 3, and 4 veins
	EARLY	LATE
Quartz	—	—
Red potassium feldspar		—
Chlorite	—	—
Pyrite	—	—
Chalcopyrite		—
Sphalerite		—
Galena		—
Tetrahedrite		—
Argentite		—
Pyrargyrite(?) or polybasite(?)		—
Carbonates		—
Native gold	—	—
Native silver		—
Telluride		—

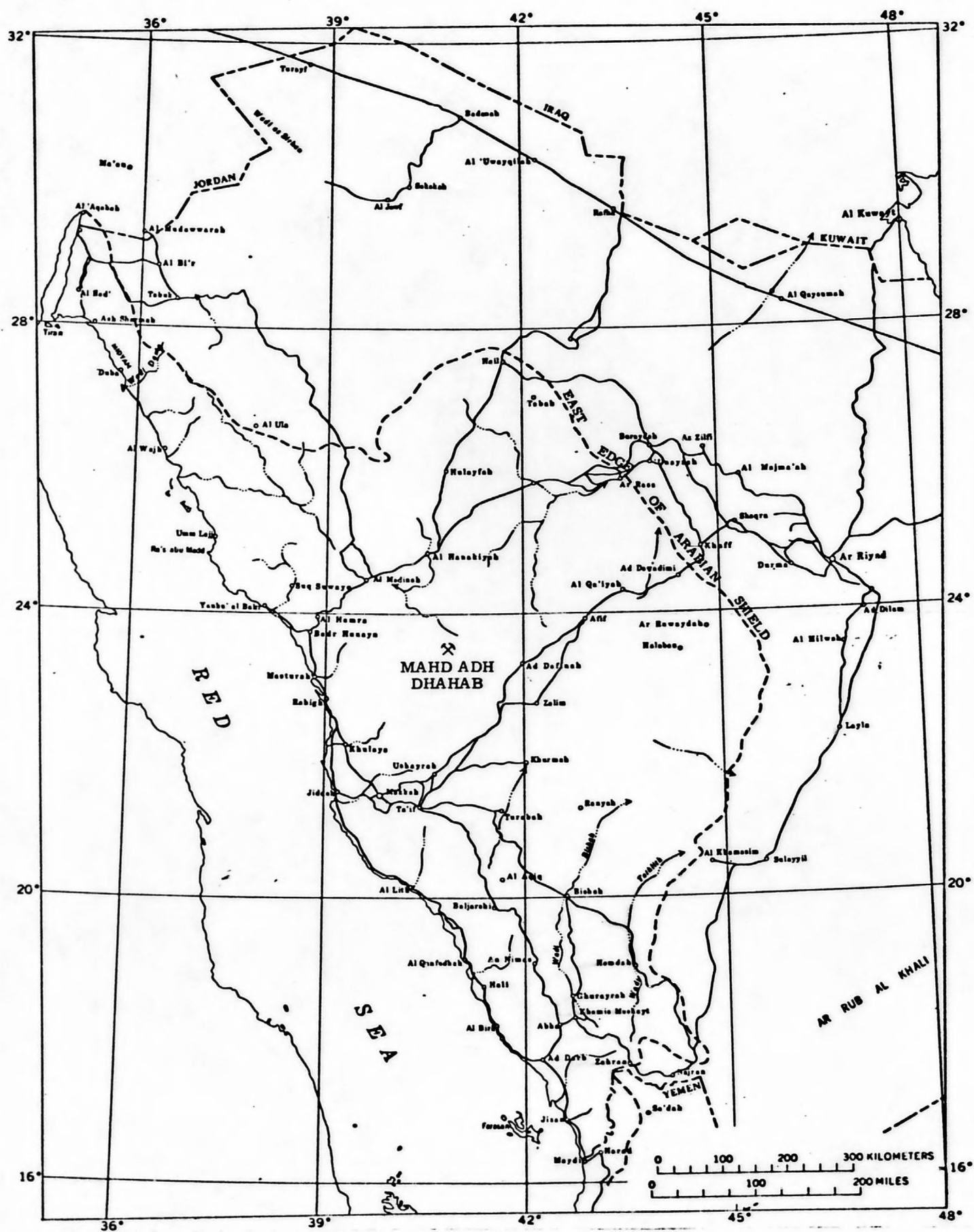
Table 2.--Sulfur isotopic values of coexisting sulfides.

Sample	Mineral	$\delta^{34}\text{S}$	T, °C	Description
107930	py	2.9	210 290	py, sl, cpy, hm, in quartz-chlorite vein
	sl	1.6		
	cpy	1.5		
107933	py	2.9	270	coarse py, cpy, and quartz
	cpy	1.4		
87376A	py	3.2		early py cubes, late sl band, quartz zones
	sl	0.8		
87375	sl	1.2		bands of quartz, chl, sl with dissem. cpy
	cpy	1.4		
69116	py	2.7		banded chl-quartz
87281	sl	2.6		sl + cpy in earlier quartz vein
69115	py	3.2	310 260	early py, sl, gn, with later banded quartz-chl-sl + cpy
	sl	1.4		
	gn	-0.7		
	sl(late)	1.6		
87281	sl	2.7		brecciated pyroclastics surrounded by quartz, cpy and sl
87283	cpy	1.6		quartz with chl stringers and cpy

Table 3.--Oxygen, carbon, and hydrogen isotopic values of quartz-carbonate pairs, microclines, chlorites, carbonates, and quartz.

Sample	$\delta^{18}O_Q$	$\delta^{18}O_C$	$\delta^{13}C_C$	δD	Description
97896	9.96	8.26	-6.67	-	coexisting vein quartz (Q) and carbonate (c).
105177	11.66	-	-	-	"
105426	11.15	11.83	-7.08	-	"
110286	8.91	9.36	-6.56	-	"
110356	8.85	19.60	-5.66	-	"
69079	10.19	15.24	-6.99	-	"
64026	10.31	-	-	-	microcline bordering quartz vein; Rb/Sr dated.
69070	9.36	-	-	-	microcline; metasomatized lower agglomerate.
69026	2.59	-	-	-65	chlorite; propylitized andesite
97668	7.22	-	-	-66	chlorite; vein selvage
69169	-	11.95	-6.46		Carbonates
69122	-	14.45	-6.27		"
64033	-	15.74	-7.43		"
69122	-	16.42	-6.48		"
64032	-	14.34	-6.41		"
64105	-	15.66	-7.58	-75	quartz; 171-278°C homog.
64120	-			-39	quartz; 151°C homog.

Figure 1.--Index map showing the location of the Mahd adh Dhahab district.



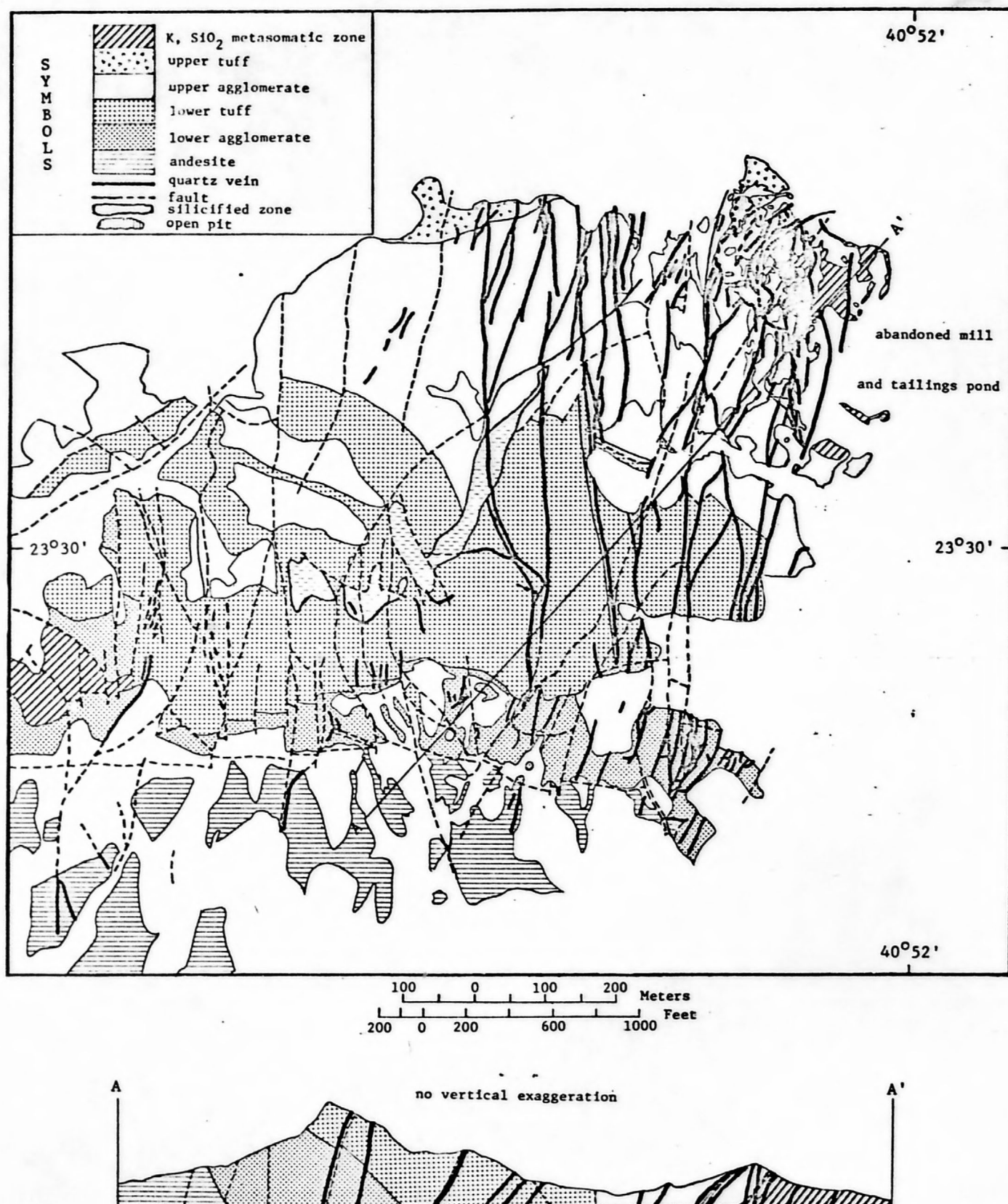
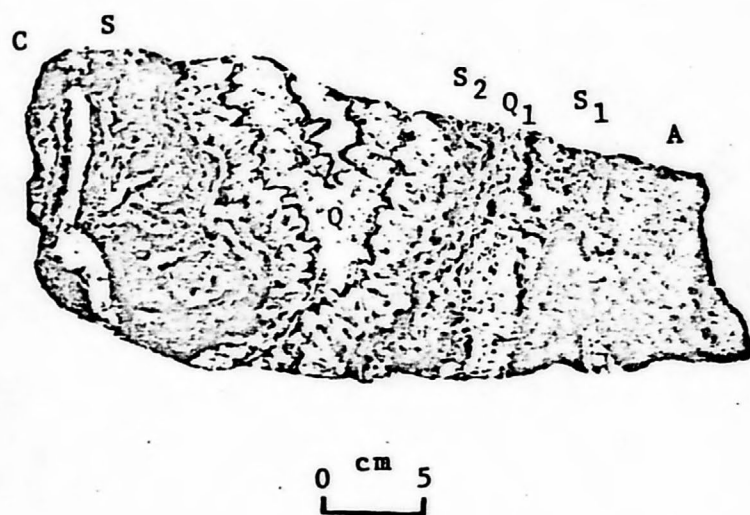


Fig. 2. Geologic Map of the Mahd adh Dhahab Deposit after Luce, Bagdady, and Roberts (1975).



	Au (ppm)	Ag (ppm)
Chloritic selvage - C	3.80	14.4
Altered tan selvage - S	46.0	50.0
Late quartz vein - Q	0.51	16.0
Altered tan selvage - S ₂	96.0	77.0
Early quartz veinlet - Q ₁	3.70	77.0
Border selvage - S ₁	0.45	17.4
Agglomerate host rock - A	0.92	7.30

Figure 3.--Type 2 quartz vein from lower agglomerate showing crustification structure. Chemical analyses are of the different banded units and were performed by atomic absorption following digestion in hydrochloric and nitric acid, at the chemical laboratory of the Saudi Arabian Directorate General of Mineral Resources, Jiddah, under the supervision of W.L. Campbell, A.Z. Khan, and Ibrahim Baraja.

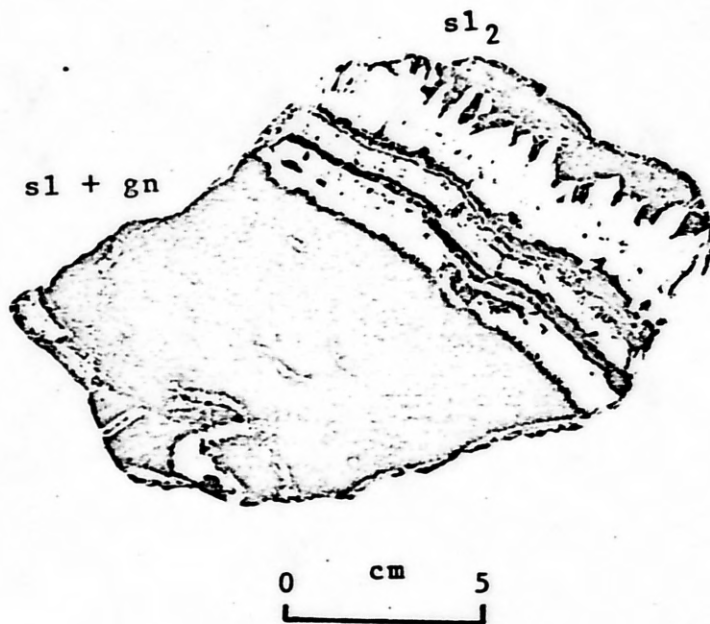


Figure 4.--Disseminated sphalerite (sl) + galena (gn) in chloritic gangue bordered by quartz veins, chlorite layers, and late-stage massive sphalerite (sl₂).