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Mineralogy, Mineral Chemistry, and Paragenesis of Gold, Silver, and Base-Metal Ores of the North Amethyst Vein System, San Juan Mountains, Mineral County, Colorado

By NORA K. FOLEY, STANTON W. CADDEY, CRAIG B. BYINGTON, and DAVID M. VARDIMAN

A study of progressive vein-related mineralization of structures in volcanic rocks of the San Luis and Bachelor calderas of the San Juan volcanic field of southwestern Colorado and its bearing on the economic potential of the area

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MINERALOGY, MINERAL CHEMISTRY, AND PARAGENESIS OF GOLD, SILVER, AND BASE-METAL ORES OF THE NORTH AMETHYST VEIN SYSTEM, SAN JUAN MOUNTAINS, MINERAL COUNTY, COLORADO

By Nora K. Foley, Stanton W. Caddey, Craig B. Byington, and David M. Vardiman

ABSTRACT

Gold-rich adularia-sericite-type mineralization occurs near the southern margin of the San Luis caldera, at the intersection of the Equity fault and the northern extension of the Amethyst fault system. Mineralized rock is confined primarily to steeply dipping structures in silicified rhyolite and dacite. Intense sericitic alteration occurs at higher levels in the vein system, and wall rock adjacent to some veins is bleached. The ores are multiply brecciated, and vein filling locally shows sedimentary textures.

Textural, mineralogical, and chemical criteria indicate that there are at least two partially coextensive associations of mineral assemblages separated by periods of brecciation and sedimentation. An older gold-bearing association consists of two fine-grained ore stages, both of which contain electrum, uytenbogaardtite, tetrahedrite, silver sulfosalts, silver sulfides and base-metal sulfides, and a manganese-rich stage containing the assemblages (1) manganese silicate + manganese carbonate minerals + quartz and (2) magnetite + hematite + pyrite + quartz. A younger crosscutting association contains calcite, adularia, fluorite, and quartz, plus the assemblages (1) coarse-grained base-metal sulfides and (2) hematite + chlorite + quartz. Quartz, manganese-rich calcite, and trace pyrite line late-stage vugs.

Mineralogic, lead-isotopic, and fluid-inclusion characteristics of the younger association are similar to those of ores of the southern and central parts of the Creede mining district. In contrast, the gold and manganese-silicate assemblages of the older association are rare to absent in the southern and central parts of the district. The local and early occurrence of the manganese and gold assemblages may indicate that they formed in a small hydrothermal cell that predated the extensive hydrothermal system from which ores of the central and southern parts of the Creede district are proposed to have been deposited (Bethke, 1988). If similar early-stage cells were present in the southern and central parts of the district, they may remain to be discovered. In the latter case, mineral assemblages that formed at early stages in the paragenesis hold the most promise for gold exploration.

INTRODUCTION

The North Amethyst vein system is located just south of the Continental Divide along the southern margin of the San Luis caldera in Mineral County, Colo. (fig. 1). The precious- and base-metal-bearing veins fill structures of the east-west-trending reverse Equity fault and steeply dipping structures related to an extension of the north-northwest-trending normal Amethyst fault system (fig. 2). These structures cut rhyolitic and rhyodacitic volcanic tuffs and flows of the central caldera cluster of the San Juan volcanic field. The veins lie just inside the northern boundary of the historic Creede mining district.

Homestake Mining Company discovered the Au-bearing ores and delineated an extensive high-grade zone adjacent to old workings of the near-surface Equity mine. Available published geologic descriptions of the mineralized rocks of the North Amethyst vein system are limited mainly to historical reports on the Equity mine (Emmons and Larsen, 1913; 1923) and some preliminary studies of the newly discovered ores (S.W. Caddey and C.B. Byington, oral commun., 1988; Plumlee and others, 1989).

This paper, part of a detailed investigation of the petrology of the complex ores of the North Amethyst vein system, concentrates on the mineralogy of the Au-rich areas exposed by underground workings. The mineralogy, mineral chemistry, and paragenesis of the ore of the North Amethyst veins, with limited petrologic interpretation, are presented here. A comparison is made with Ag- and base-metal-rich mineralization of the southern Amethyst, Bulldog Mountain, P, and OH veins of the main Creede mining district. Only a limited amount of petrologic interpretation is included in this paper; instead, the paper focuses on the mineralogy and stage assemblages. The petrology, fluid-inclusion thermometry, and lead-isotopic geochemistry of the vein system are described by Foley (1990).

ACKNOWLEDGMENTS

Homestake Mining Company discovered the veins that constitute the North Amethyst system in 1983 as the
result of an extensive drilling program (S.W. Caddey and C.B. Byington, oral commun., 1988; Plumlee and others, 1989). Homestake Mining Company generously provided access to exploratory workings of the North Amethyst vein and the opportunity to study some aspects of the newly discovered mineralization. Stanton W. Caddey, Craig B. Byington, and David M. Vardiman, Homestake Mining Company geologists who discovered and developed the veins, contributed their time and their knowledge of the underground workings. They provided access to samples collected throughout the development of the underground workings and therefore improved the interpretations of hand-sample textures and vein relations.

The senior author would like to thank Paul B. Barton, Jr., and Philip M. Bethke of the U.S. Geological Survey (USGS) for suggesting that she undertake this project and for providing an initial set of samples. Geoffrey S. Plumlee of the USGS also contributed samples and took some of the underground photographs. Careful reviews of this manuscript by Robert A. Ayuso (USGS), Paul B. Barton, Jr., Philip M. Bethke, James R. Craig (Virginia Polytechnic Institute and State University), Geoffrey S. Plumlee, and Gilpin R. Robinson, Jr. (USGS), have improved both its content and presentation. Much of this manuscript constitutes a part of the senior author’s Ph.D. dissertation. Conversations with H.E. Belkin, J.M. Hammarstrom, D.O. Hayba, Pamela Heald, J.S. Huebner, M.J.K. Flohr, P.M. Okida, J.F. Slack, and D.S. Sweetkind, all colleagues at the USGS, were invaluable throughout the duration of the study. J.J. McGee and J.V. Emery provided guidance in use of the electron microprobe.
GEOLOGIC SETTING

The San Juan volcanic field is an erosional remnant of a much larger volcanic field that covered part of the southern Rocky Mountains in the middle Tertiary (Steven, 1975; Lipman and others, 1978). The present extent of the major ash-flow tuffs and their associated calderas is shown in figure 1. The volcanic field rests upon an eroded base consisting of crystalline metavolcanic, metasedimentary, and granitoid rocks of Proterozoic age, and, possibly, remnants of Paleozoic, Mesozoic, and Tertiary sedimentary rocks (summarized by Steven and Lipman, 1976; Lipman and others, 1978; Baars and Stevenson, 1984; Gries, 1985; Bickford, 1988). The composite volcanic field consisted dominantly of andesite-dacite lavas and associated volcaniclastic rocks erupted from precaldera-stage volcanoes in early Oligocene time, mainly between 35 and 30 Ma. A large negative Bouguer gravity anomaly underlies much of the San Juan volcanic field and is believed to reflect a large composite batholith emplaced beneath the field (Plouff and Pakister, 1972). From about 28 to 26 Ma (table 1), the central San Juan volcanic field experienced widespread silicic ash-flow eruptions and related caldera subsidence (Lipman and others, 1970, 1978; Lipman and Sawyer, 1988; Lanphere, 1988). The mineral deposits of the Creede district are located in a complex set of nested calderas formed by the pyroclastic eruptions.

The most volumetrically important sediments that accumulated within the San Juan area during the middle...
The complex history of movement and reactivation along faults of the Creede graben has been described in detail by Steven and Ratté (1965). Normal faults are the dominant structural element controlling the location of mineralized rock in the central San Juan area, and most are related to evolution of the Oligocene calderas (fig. 2). The reverse Equity fault, which formed by fault-bounded uplift of a triangular block presumably related to a shallow underlying intrusion (Lipman and Sawyer, 1988), is also mineralized. The dominant structural features are the well-formed caldera expressions and the north-northwest-trending Creede graben, which initially formed as a keystone graben in the Bachelor caldera (fig. 1). The major faults from west to east include the Alpha-Corsair, Bulldog Mountain, Amethyst, Midwest-Knoess, and Solomon-Holy Moses faults. The OH, P, and other minor faults cut the central block of the graben, are nearly vertical, and had primarily strike-slip motion (S.W. Caddey, written commun., 1985).

**AGE OF MINERALIZATION**

A single radiometric age determination (table 1) on adularia (40Ar/39Ar incremental heating ages) from drill-core from the North Amethyst vein system indicates an age of 25.1±0.1 Ma (M.A. Lanphere, written commun., 1987). The adularia is found with base-metal minerals of a late-stage hydrothermal event that occurred along the North Amethyst vein system. The Au-bearing ores predate this base-metal and vein adularia stage (Foley and Vardiman, 1988), so their age falls in the interval between that of the youngest host rock, which is the Nelson Mountain Tuff, at 25.1 Ma, and the vein adularia, at 25.1 Ma. At least five volcanic units were deposited as part of the San Luis caldera complex after deposition of the oldest host rock, which is the Nelson Mountain Tuff, at 26.1 Ma, and the vein adularia, at 25.1 Ma.
Figure 3.—Plan map showing the levels of the North Amethyst vein system along the northern Amethyst and Equity faults. Sample localities are shown by solid dots; localities off the vein are on the decline.

PREVIOUS STUDIES

Mineralized rock in the central part of the San Juan area has been the subject of numerous reports and papers since the area was first mined in the early 1880’s. The relationship of the mineralization to regional geologic setting was first described in detail by Steven and Ratté (1960, 1965) and Steven (1968). The area in and around the Creede mining district has been the subject of continuing research since the late 1950’s by the USGS and by several universities. Their work has been published in a series of papers on the Creede district (Steven and Eaton, 1975; Bethke and others, 1976; Roedder, 1977; Barton and others, 1977; Wetlaufer, 1977; Bethke and Rye, 1979; Hayba and others, 1985; Plumlee, 1989; Foley, 1990; Foley and others, 1990). Historic accounts of ores of the Equity mine and a summary of preliminary reports of the more recently discovered North Amethyst ores are described in Foley (1990).

STYLE OF MINERALIZATION

The vertical and lateral extent, and overall shape, of ore occurrences of the North Amethyst vein system (fig. 3) is not well delineated because of the complex nature of the mineralized rock and the complex structural character of the veins. A detailed structural analysis of the North Amethyst vein system is necessary to resolve composite vein and structure relations. Some general observations, however, can be made relative to ore zones of the Bulldog and Amethyst vein systems of the main mining district at Creede as described by Plumlee (1989) and to ore zones of the OH and P veins.
Table 2.—Stages of mineralization and brecciation events of the North Amethyst vein system

<table>
<thead>
<tr>
<th>Stage</th>
<th>North Amethyst vein mineral assemblage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Base-metal–silica association:</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Mn-calcite » quartz &gt; pyrite</td>
</tr>
<tr>
<td>Breccia-4</td>
<td>Mn-calcite cement, fragments of earlier assemblages, breccias and minerals [wall rock]</td>
</tr>
<tr>
<td>2b</td>
<td>Quartz &gt; calcite &gt; rhodocrosite &gt; adularia &gt; fluorite &gt; chlorite = pyrite</td>
</tr>
<tr>
<td>Breccia-3</td>
<td>Quartz + chlorite cement, fragments of wall rock, and coarse stage-1 sulfides</td>
</tr>
<tr>
<td>1b</td>
<td>Quartz &gt; sphalerite &gt; galena &gt; pyrite &gt; chalcopyrite &gt; tetrahedrite</td>
</tr>
<tr>
<td>1a</td>
<td>Quartz &gt; pyrite = hematite &gt; chlorite</td>
</tr>
<tr>
<td>Breccia-2</td>
<td>Quartz cement, fragments of stages alpha and beta, wall rock, sedimented structures, black quartz breccia</td>
</tr>
<tr>
<td>Mn-Au association:</td>
<td></td>
</tr>
<tr>
<td>Beta-2</td>
<td>Sphalerite &gt; galena &gt; chalcopyrite &gt; pyrite &gt; tetrahedrite &gt; Ag- and Au-bearing sulfide minerals &gt; electrum [magnetite, hematite]</td>
</tr>
<tr>
<td>Breccia-1</td>
<td>Quartz cement, fragments of stages alpha and beta</td>
</tr>
<tr>
<td>Beta-1</td>
<td>Sphalerite &gt; galena &gt; chalcopyrite &gt; pyrite &gt; tetrahedrite &gt; Ag- and Au-bearing sulfide minerals &gt; electrum [magnetite, hematite]</td>
</tr>
<tr>
<td>Alpha</td>
<td>Rhodocrosite = quartz &gt; Mn-calcite = rhodonite &gt; K-feldspar &gt;&gt; sphalerite &gt; pyrite &gt; galena &gt; [barite]</td>
</tr>
</tbody>
</table>

Mineralized rock of the North Amethyst vein system fills structures related to the northern Amethyst and Equity faults. The North Amethyst veins are mineralogically complex, and the assemblages can be divided into two contrasting associations: a Mn-Au association and a base-metal–silica association. The minerals found in each association are grouped in table 2 according to the stages in which they are typically found and are discussed in greater detail herein. Both mineral associations occur along almost the entire extent of the underground workings, a distance of roughly 600 m along strike of the vein. The base-metal association is displayed over a vertical extent of about 150 m, whereas minerals of the Mn-Au association occur over an apparently larger vertical interval of 600 m.

Character of the ore zones

Ore zones of the main mining district typically are of much greater horizontal than vertical extent. For example, the ore zone comprising the Bulldog Mountain vein system was mined for approximately 3 km laterally and 200 to 300 m vertically (Plumlee, 1989). The combined productive ore zones of the central and southern Ame-
nated here as stage alpha, was deposited primarily in open fractures, in some cases around preexisting barite and quartz crystals. The cryptocrystalline material shows a faint but clearly visible mineralogic banding on aged surfaces that is due to manganese oxidation. Stage alpha also shows some evidence for replacement of volcanic wall rock. Embayed remnant feldspar grains are caught up in a fine-grained “frozen mush” consisting dominantly of rhodocrosite, manganese pyroxenoids and other carbonate minerals, and quartz. The second stage of the Mn-Au association, stage beta, consists of fine-grained precious- and base-metal-sulfide-rich seams that cut and recrystallized the stage-alpha minerals.

The later, base-metal-silica association minerals filled open spaces in fractures and breccias (fig. 5). Although many veins filled completely, some large vugs lined with crystals of the latest stages remained open. Minerals of this association are generally coarser grained than those of the Mn-Au association and show crustification sequences. Dissolution and replacement textures are also common.

VEIN MINERALOGY AND COMPOSITIONAL RANGES

Detailed descriptions of ore and gangue minerals are given here for common or economically important minerals and for those used to constrain conditions of ore formation. The minerals are listed in alphabetical order and grouped by type for convenience; sample locations are shown in figure 6. Table 3 describes the relative abundance and distribution of minerals in each assemblage and includes all minerals reported for the deposit. Compositional ranges for carbonate minerals, electrum, manganese silicates, and sphalerites described in table 4 are listed in tables 5 through 8. Sample distribution, selection method, analytical details, and analyses are described by Foley (1990). Ore minerals were identified on the basis of descriptions presented by Ramdohr (1969) and Craig and Vaughan (1981).

Ag-SULFIDE AND Ag-Cu-SULFIDE MINERALS

Four Ag- and Cu-bearing sulfide minerals occur in the Au-rich stage of mineralization: acanthite, jalpaite, mckinstryite, and, possibly, stromeyerite. These phases were identified on the basis of optical properties and analysis by an energy dispersive system but were not examined by X-ray diffraction. Where possible, the compositions were estimated by energy dispersive analysis and electron microprobe analysis. The Cu-Ag-S minerals do not yield reliable temperatures of specific conditions of ore deposition because of their exceedingly fast reequilibration rates (Skinner, 1966) but are included in this discussion because of their economic importance.

Acanthite (Ag₂S) - Argentite (Ag₂S).—Small (<0.03 mm) irregularly shaped grains of acanthite occur alone in quartz gangue and at the margins of galena, pyrrargyrite, and chalcopyrite grains and as inclusions in pyrite and galena. The Ag sulfide is contemporaneous with chalcopyrite and pyrrargyrite. It is also found both alone and with polybasite, in association with galena, with chalcopyrite and electrum, and rimming and enveloping rounded grains of galena. Some small (<0.01 mm), rounded grains are contained within galena; however, no clear-cut exsolution textures were observed.

Jalpaite (Ag₅CuS₆).—Soft, light-gray, Ag- and Cu-bearing sulfide mineral grains, 10 to 25 μm in width, are interfingered with or are enclosed by acanthite that is in turn bordered by chalcopyrite and galena. This mineral is tentatively identified as jalpaite on the basis of an absence of Sb or As, high Ag relative to Cu, good polish, and distinct anisotropism without the striking blue-violet tints of stromeyerite. The mineral also occurs in aggregates with acanthite and Ag-bearing tetraedrite that are enclosed by gangue.

Mckinstryite (Ag₁₃Cu₀₇S).—Anhedral crystals of mckinstryite (15 μm wide) are interfingered locally with acanthite, jalpaite, uytenbogaardtite, and, possibly, stromeyerite. The mineral also occurs as small <20-μm-wide anhedral grains in aggregates interfingered with chalcopyrite and acanthite, and with wire silver the mckinstryite fills cavities and microfractures that cut sulfide-rich zones. Although optical properties of this phase closely resemble those of stromeyerite, the two Cu-Ag sulfides can be distinguished optically when interfingered. Identification was confirmed by electron microprobe where possible.

BARITE

Barite (BaSO₄) was not found in samples collected from underground workings. However, tabular molds up to 3 cm in length resembling the crystal habit of barite do occur in samples from the lower elevations in the workings, especially to the south. The mineral is found in drillcore samples from the southern end of the underground workings and in core drilled to the south of the exploratory workings. Tabular and bladed crystals of colorless to pale-cream barite ranging in length from 1 to 10 mm occur in veinlets with quartz, calcite, and minor pyrite.

BOURNONITE

A gray-white phase (PbCuSbS₃) occurs with galena rimming corroded sphalerite and as small aggregates of
Metal-silica association is colorless to pale yellow and mainly contains MnCO₃. The range in manganese contents from about 10 to about 30 mole percent MnCO₃.

The fine-grained nature of these samples hindered their characterization; hence, rhodocrosite, and quartz. Calcite of this stage shows a fine grained (crystals <50 μm in width) and is intergrown with microcrystalline Mn-silicates, kutnahorite, and a calcian rhodocrosite. Each mineral has a range of compositions. Representative analyses are given in table 5; all analyses are plotted in figures 7 and 8.

Calcite (Ca₀.₇₅₋₀.₉₅Mn₀.₂₆₋₀.₆₉CO₃).—Three calcites are distinguished on the basis of paragenetic position, color, and grain size; however, they overlap in part in composition. Calcite associated with the Mn-Au stages is very fine grained (crystals <50 μm in width) and is intergrown with microcrystalline Mn-silicates, kutnahorite, rhodocrosite, and quartz. Calcite of this stage shows a range in manganese contents from about 10 to about 30 mole percent MnCO₃.

Calcite that occurs in the early stages of the base-metal-silica association is colorless to pale yellow and generally has lower manganese contents than other analyzed carbonate minerals (9 to 15 mole percent MnCO₃). The crystals range in size from a few millimeters to 2 to 3 cm and are intergrown with 0.1- to 1.0-mm-wide grains of oligonite, the manganese siderite. The calcite contains virtually no magnesium and generally less than 1 mole percent FeCO₃.

The last stage of the base-metal-silica association is composed dominantly of a pink to pale-cream manganese calcite, especially in the northern part of the ore zone. Thick crusts of crystalline carbonate, up to 15 cm in thickness, are found mainly lining late-stage vugs and open spaces in the vein. The crusts are dominantly pale cream in color and contain thin bands of darker tan to rust-colored carbonate. The compositional range for this calcite is similar to calcite of the Mn-Au stages.

Kutnahorite (Ca₀.₄₇Mn₀.₅₃CO₃).—A Mn- and Ca-bearing carbonate mineral having a composition equivalent to kutnahorite, the manganese analog of dolomite, has been identified intergrown with fine-grained rhodocrosite, Mn-calcite, and Mn-silicates of the Mn-Au association (table 5). X-ray diffraction analyses of the mixture suggest primarily quartz, rhodonite, pyroxmangite, and carbonate minerals having a wide range of Ca-Mn solid solution. Microprobe analyses of the carbonate minerals yield some compositions in the range Ca₀.₇₀Re₀.₃₄ to Ca₀.₅₃Re₀.₄₇ with less than 1 percent (Fe,Mg)CO₃. The material is too fine grained to permit single-crystal X-ray study to determine if the kutnahorite is ordered; long-range cation ordering of kutnahorite is not detectable by conventional powder X-ray diffraction techniques (Pecor and others, 1987).

Rhodocrosite (Mn₀.₆₆₋₀.₇₆Ca₁₃₋₁₉CO₃).—In the Mn-Au association, rhodocrosite is the most abundant carbonate mineral and occurs in a microcrystalline (≤50 μm) admixture composed mainly of rhodonite, quartz, and Mn-calcite. The admixture is zoned mineralogically on the scale of hundreds of micrometers. Coarser rhodocrosite (up to 0.5 mm in width) occurs in some of the zones and in patches that cut across zones; the patches consist of mats of intergrown curls that resemble poole hair. Compositions of the poole-hair rhodocrosites are similar to, and as calcic as, those of the finer grained material (up to Ca₀.₅₃Re₀.₅₇).

In the base-metal-silica association, rhodocrosite is associated with amethystine quartz and fluorite. Rhodocrosite also occurs in most later stages of the North Amethyst paragenesis in small (<5 mm wide) veinlets, and as 1- to 5-mm-wide, near-gem-quality crystals that line small vugs and are perched on minerals from earlier stages.

Siderite, manganese. —Fe- and Mn-bearing oligonite (Fe₄.₅₄Mn₃.₃₂Ca₆.₁₂Mg₄.₂₀CO₃) is found in one assemblage of the base-metal-silica association. The mineral occurs as discrete crystals 0.1 to 1.0 mm wide within a coarser calcite. The carbonate minerals are intergrown primarily

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4 Capobianco and Navrotzky (1987) recommend that the name kutnahorite be applied to the 50:50 phase having dolomite-type ordering and that naturally occurring phases lacking dolomite-type ordering be termed pseudokutnahorite. The fine-grained nature of these samples hindered their characterization; hence, ordering (or the lack of ordering) is not implied by the use of the name kutnahorite.
ORES OF THE NORTH AMETHYST VEIN SYSTEM, COLORADO
with chlorite, quartz, hematite, and minor adularia and sphalerite and were deposited just after a major period of base-metal mineralization.

Wetlaufer (1977), in a study of carbonate minerals occurring in the southern part of the Creede district, documented significant compositional variations in rhodocrosite and siderite. She found that Bulldog Mountain rhodocrosite compositions from the early pre-ore stage lie along the FeCO$_3$-MnCO$_3$ join and typically contain 68 to 93 mole percent MnCO$_3$. CaCO$_3$ contents vary from 2 to 15 mole percent, and MgCO$_3$ contents are generally less than 4 mole percent. A later intra-ore siderite-manganosiderite stage was found to contain 33 to 94 mole percent FeCO$_3$, 6 to 67 mole percent MnCO$_3$, 1 to 13 mole percent CaCO$_3$, and 1 to 10 mole percent MgCO$_3$.

**CHALCOPYRITE**

Chalcopyrite (CuFeS$_2$) is abundant in three sulfide-bearing stages of mineralization and is not found in stages composed dominantly of gangue. Size is highly variable; diameters of the irregular grains range from a few micrometers to a few millimeters. Earlier stage occurrences are generally finer grained than later stage occurrences.

Chalcopyrite is generally formed later than and replaces sphalerite. Extensive chalcopyrite disease textures (Barton and Bethke, 1987) are present in all sphalerite-bearing stages. The disease consists of
Table 3.—Assemblages and minerals of the North Amethyst vein system

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<tr>
<th>Mineral</th>
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<th>Alpha</th>
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<th>Beta 2</th>
<th>1-a</th>
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<td>x</td>
<td>x</td>
<td>-</td>
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</table>

micrometer-sized chalcopyrite grains that are rounded to rod shaped; they are generally concentrated at the rims of sphalerite crystals.

In earlier stages, chalcopyrite is found in sphalerite and as somewhat larger separate grains bordering and corroding sphalerite. Xenomorphic grains of chalcopyrite are intergrown with galena, tetrahedrite, silver minerals, and electrum, and together these minerals fill interstices around crystals of sphalerite and pyrite (fig. 4E and F).

In the later stages, chalcopyrite occurs as anhedral grains bordering sphalerite and as common intergrowths with galena. It also occurs as isolated grains that are entirely surrounded by quartz and carbonate gangue.

Oxidation of the chalcopyrite to chalcocite and covellite has been reported for some samples of the North Amethyst ores (S.W. Caddey and D.M. Vardiman, oral commun., 1988); this type of oxidation was not observed in any of the samples examined in this study.

**CHLORITE**

In the North Amethyst veins, chlorite \((A_5-Z_4)O_{10}(OH)_8\), where \(A=Al,Fe^{2+},Fe^{3+},Mg,Mn\) and \(Z=Al,Fe^{3+},Si\) occurs primarily in the later, base-metal-silica association where it is intergrown with hematite, quartz, and pyrite. Assemblages containing the green, fine-grained (<100-μm-wide flakes) layer silicate formed just prior to, and concurrently with, the period of major base-metal-sulfide mineralization in the later association. Some chlorite is also intergrown with quartz and calcite deposited after the base-metal sulfides.
VEIN MINERALOGY AND COMPOSITIONAL RANGES

Table 4.—Locations and descriptions of samples from the North Amethyst vein system analyzed by electron microprobe

<table>
<thead>
<tr>
<th>Sample no.</th>
<th>Location</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>DMV-1</td>
<td>10250 level, E-126+75’N</td>
<td>Coarse-grained (up to 1 cm wide) stage-1 lemon-yellow crystals of sphalerite intergrown with galena and chalcopyrite in veinlets 2 mm to 1 cm wide cutting bleached volcanic rock.</td>
</tr>
<tr>
<td>DMV-3</td>
<td>10325 level, 2’ south</td>
<td>Seam of fine-grained electrum-bearing sulfide at contact of volcanic wall rock with breccia of stage beta.</td>
</tr>
<tr>
<td>GSP-MM-10-86</td>
<td>10250 level, E-125+10’N</td>
<td>20-cm-wide seam of gold-bearing fine-grained sulfides of stage beta in fracture cutting vein of stage alpha. Rounded grains of sphalerite intergrown with chalcopyrite, galena, electrum, and other precious- and base-metal minerals of stage beta.</td>
</tr>
<tr>
<td>GSP-MM-10-86</td>
<td>10250 level, E-125+35’N</td>
<td>Large (up to 1 cm wide) color-banded crystals of sphalerite of stage 1 intergrown with quartz and chlorite, coated by stage-3 pink calcite.</td>
</tr>
<tr>
<td>GSP-MO-2-86</td>
<td>10250 level, E-126+50’N, in scram off vein.</td>
<td>Coarse veins of honey-brown sphalerite, galena, chalcopyrite, pyrite, and quartz of stage 1 cutting a meter-wide brecciated vein of Mn-carbonate, Mn-silicate, and quartz.</td>
</tr>
<tr>
<td>NKF-A-1-87</td>
<td>10250 level, E-127+140’N</td>
<td>Coarse- to medium-grained sulfides of stage 1 cutting a meter-wide brecciated vein of Mn-carbonate, Mn-silicate, and quartz.</td>
</tr>
<tr>
<td>NKF-K-11-87</td>
<td>10250 level, E-85</td>
<td>Fluorite at base overlain by thick crust of calcite color banded from pink to pale-cream.</td>
</tr>
<tr>
<td>NKF-M-13-87</td>
<td>10250 level, E-120</td>
<td>Small (1 to 2 mm wide) rounded sphalerite grains of stage alpha rimmed with chalcopyrite disease, contained in fine-grained admixture of rhodonite, rhodochrosite, and quartz.</td>
</tr>
<tr>
<td>NKF-Q-17-87</td>
<td>10250 level, 25 ft from north face</td>
<td>Meter-wide vein of Mn-carbonates and Mn-silicates containing disseminated galena (stage alpha), sulfides, quartz, and rhodochrosite.</td>
</tr>
<tr>
<td>NKF-R-18-87</td>
<td>10250 level, E-126+25’S</td>
<td>Banded hematite, magnetite, sulfides, quartz, and electrum (stage alpha). Cut by fractures containing coarse-grained sulfides of stage 1.</td>
</tr>
<tr>
<td>NKF-X-24-87</td>
<td>10250 level, 25 ft from north face</td>
<td>Fine-grained carbonate, Mn-silicate, and quartz material of stage alpha. Disseminated sulfides in recrystallized Mn-carbonates and Mn-silicates. Fine-grained minerals of stage alpha cut by veins containing red-brown to yellow sphalerite of stage 1, pyrite, chalcopyrite, and quartz and calcite gangue.</td>
</tr>
<tr>
<td>NKF-BM-65-87</td>
<td>10250 level, S2 raise</td>
<td>Rounded cobble of fine-grained sulfide containing gold and sphalerite (stage beta) in breccia.</td>
</tr>
</tbody>
</table>

No quantitative compositional data are available for North Amethyst chlorites. They are iron rich like other Creede chlorites (Barton and others, 1977) and contain substantial manganese (by energy dispersive analysis).

Electrum (Au$_{0.3-0.5}$Ag$_{0.7-0.3}$) is characteristic of the early, Mn-Au association of the North Amethyst paragenesis. The mineral is intricately intergrown with a variety of base- and precious-metal sulfides and generally formed late with respect to the base-metal sulfides (fig. 4F). The whitish-yellow Ag-rich grains occur most frequently as small rounded inclusions (3 to 25 μm) in chalcopyrite and uytenbogaardtite, although they are also found in acanthite, pyrargyrite, and galena. More rarely, electrum occurs as small (3 to 30 μm) separate grains in quartz gangue and occurs adjacent to sphalerite aggregates rimmed with galena and chalcopyrite. Larger grains, up to 75 μm in diameter and visible in hand specimen, occur with pyrargyrite, chalcopyrite, and galena. They fill interstices between, and in part replace, sphalerite grains at lower elevations in the mine. Representative electrum compositions are given in table 6 and plotted in figure 9. Analyzed electrum contained entirely within pyrite (Nos. 6, 7, table 6) has slightly higher Au...
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TABLE 6.—Representative electron microprobe analyses of carbonate minerals from the North Amethyst vein system
[Values in weight percent; collected by: NKF, Nora K. Foley; GSP, Geoffrey S. Plumlee]

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Atomic proportion

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<tbody>
<tr>
<td>Ag⁺¹</td>
<td>.72</td>
<td>.65</td>
<td>.58</td>
<td>.52</td>
<td>.54</td>
<td>.49</td>
<td>.48</td>
</tr>
<tr>
<td>Au⁺²</td>
<td>.28</td>
<td>.35</td>
<td>.42</td>
<td>.48</td>
<td>.46</td>
<td>.51</td>
<td>.52</td>
</tr>
</tbody>
</table>

1. NKF-M-13-87, fine-grained pale-pink calcite from stage alpha-1.
2. NKF-BB-54-87, fine-grained "kutnahorite" from stage alpha-2.
3. NKF-BB-54-87, medium-grained "poodle-hair" rhodocrosite from stage alpha-2.
4. NKF-BB-54-87, medium-grained pink rhodocrosite of stage alpha-1.
5. GSP-MN-24-87, colorless to pale-cream calcite of stage 2 (intergrown with no. 7).
6. GSP-MN-1-87, yellowish manganese siderite of stage 2 (intergrown with no. 6).
7. NKF-K-11-87, true pink manganese calcite of stage 3.

Table 6.—Representative electron microprobe analyses of electrum from the North Amethyst vein system
[Values in weight percent; collected by: NKF, Nora K. Foley; GSP, Geoffrey S. Plumlee]

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<tr>
<td>Ag</td>
<td>57.52</td>
<td>49.33</td>
<td>42.77</td>
<td>36.76</td>
<td>38.80</td>
<td>33.40</td>
<td>32.90</td>
</tr>
<tr>
<td>Au</td>
<td>40.74</td>
<td>49.31</td>
<td>56.02</td>
<td>61.31</td>
<td>59.65</td>
<td>64.34</td>
<td>65.73</td>
</tr>
<tr>
<td>Total</td>
<td>98.26</td>
<td>98.64</td>
<td>98.79</td>
<td>98.07</td>
<td>98.45</td>
<td>97.74</td>
<td>98.63</td>
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Atomic proportion

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<tbody>
<tr>
<td>Ag⁺¹</td>
<td>.72</td>
<td>.65</td>
<td>.58</td>
<td>.52</td>
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<td>.28</td>
<td>.35</td>
<td>.42</td>
<td>.48</td>
<td>.46</td>
<td>.51</td>
<td>.52</td>
</tr>
</tbody>
</table>

1. NKF-AA-27-87, silver-rich electrum grain contained in matrix of siliceous quartz.
2. NKF-AA-27-87, electrum contained in chalcopyrite.
3. NKF-AA-27-87, electrum rimmed by acanthite and uytenbogaardtite.
4. NKF-BM-65-87, electrum grain intergrown with sphalerite, galena, and chalcopyrite.
5. NKF-BM-65-87, electrum rimmed by silver minerals including acanthite and uytenbogaardtite.
6. GSP-MM-10a, gold-rich electrum grain entirely rimmed by pyrite.
7. DMV-3, gold-rich electrum grain contained in pyrite.
TABLE 7.—Representative electron microprobe analyses of Mn-silicate minerals from the North Amethyst vein system
([Values in weight percent; -, not detected; collected by: NKF, Nora K. Foley; GSP, Geoffrey S. Plumlee; EQ, David M. Vardiman])

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<tbody>
<tr>
<td>SiO₂</td>
<td>45.41</td>
<td>45.64</td>
<td>45.35</td>
<td>47.10</td>
<td>47.06</td>
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<tr>
<td>Al₂O₃</td>
<td>0.13</td>
<td>0.27</td>
<td>0.13</td>
<td>0.55</td>
<td>0.20</td>
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<tr>
<td>Fe₂O₃</td>
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<td>5.87</td>
<td>2.70</td>
<td>2.21</td>
<td>2.98</td>
<td>1.03</td>
<td>1.73</td>
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<tr>
<td>FeO</td>
<td>47.95</td>
<td>47.01</td>
<td>45.22</td>
<td>43.23</td>
<td>41.79</td>
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<tr>
<td>MnO</td>
<td>0.42</td>
<td>0.31</td>
<td>0.56</td>
<td>0.39</td>
<td>0.71</td>
<td>0.03</td>
<td>0.47</td>
</tr>
<tr>
<td>MgO</td>
<td>3.65</td>
<td>1.51</td>
<td>1.73</td>
<td>6.02</td>
<td>6.32</td>
<td>10.59</td>
<td>6.78</td>
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<tr>
<td>CaO</td>
<td>99.47</td>
<td>100.61</td>
<td>98.26</td>
<td>101.49</td>
<td>100.50</td>
<td>100.18</td>
<td>91.78</td>
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</table>

Number of ions on the basis of 6 or 36 oxygen

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<tbody>
<tr>
<td>Si IV</td>
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<td>1.98</td>
<td>2.00</td>
<td>1.99</td>
<td>2.00</td>
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<td>T site</td>
<td>1.98</td>
<td>1.99</td>
<td>2.00</td>
<td>2.00</td>
<td>2.00</td>
<td>1.99</td>
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<tr>
<td>Al VI</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.01</td>
<td>0.01</td>
<td>-</td>
</tr>
<tr>
<td>Fe³⁺</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.00</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Fe²⁺</td>
<td>0.07</td>
<td>0.21</td>
<td>0.10</td>
<td>0.08</td>
<td>0.11</td>
<td>0.04</td>
</tr>
<tr>
<td>Mn²⁺</td>
<td>1.77</td>
<td>1.72</td>
<td>1.78</td>
<td>1.61</td>
<td>1.55</td>
<td>1.51</td>
</tr>
<tr>
<td>Mg</td>
<td>0.08</td>
<td>0.02</td>
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</tr>
<tr>
<td>Ca</td>
<td>0.17</td>
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<td>0.08</td>
<td>0.27</td>
<td>0.29</td>
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<tr>
<td>M₁,M₂</td>
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<td>2.03</td>
<td>2.00</td>
<td>2.00</td>
<td>2.00</td>
<td>2.03</td>
</tr>
</tbody>
</table>

4. EQ–16–1, fine crystals of rhodonite intermixed with rhodocrosite, calcite, and quartz.
5. NKF–Q–17–87, colorless to pink crystals of rhodonite adjacent to vuggy quartz.
7. Inesite from the Equity vein, Equity mine; FeO determined colorimetrically, MnO by XRF, and all other elements by gravimetrics (Van Loenen, 1980).

FLUORITE

Large cubes of fluorite (CaF₂) up to 13 mm on a side are found lining vugs, veins, and cavities in the North Amethyst vein system. The crystals are most frequently green; purple and colorless varieties are less common. No clear-cut relation between color and paragenetic position was discerned. Fluorite generally forms late in the paragenesis and usually grows on quartz; however, the fact that the bases of many late-stage quartz crystals contain numerous small octahedra of colorless to pale-green fluorite indicates near-contemporaneous deposition. Fluorite crystals are sometimes dusted with fine-gray quartz sediment that is overgrown by coarse cream-to-pink, manganese-rich calcite and (or) pyrite (fig. 5F). In small vugs, fluorite may follow quartz and adularia.

GALena

Galena (PbS) occurs in both of the principal sulfide stages of the North Amethyst paragenesis. It generally is intergrown with chalcopyrite and is similar to chalcopyrite in size and texture where they are found together.

In the Mn-Au association, galena is found as both isolated anhedral grains and aggregates bordering and corroding sphalerite and pyrite in a gangue of fine-grained Mn-carbonate minerals, Mn-silicates, and quartz. In sulfide-rich assemblages, galena is usually in mutual contact with chalcopyrite, pyrargyrite, acenthite, uytenboogaardtite, and polybasite. Bouronite rims are present on some galena. Micrometer-wide rounded grains of galena occur along fractures in the outer zones of sphalerite crystals, and larger grains (10 to 100 µm wide) occur in an almost equigranular mix of sphalerite and chalcopyrite.

In the base-metal-mineral association, anhedral grains and aggregates of 2- to 3-mm-wide galena grains are found bordering sphalerite or enclosed by gangue. The galena is often intergrown with chalcopyrite or tetrahedrite. The generally xenomorphic texture of galena and the occurrence along cracks in sphalerite crystals suggest that it may have formed later than did sphalerite and pyrite. Skeletal crystals of galena also grow on sphalerite and project into quartz- and calcite-filled vugs.
ORES OF THE NORTH AMETHYST VEIN SYSTEM, COLORADO

Table 8.—Representative electron microprobe analyses of sphalerite from the North Amethyst vein system

[Values in weight percent; collected by: NKF, Nora K. Foley; GSP, Geoffrey S. Plumlee]

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<td>0.80</td>
<td>0.60</td>
<td>1.58</td>
<td>0.67</td>
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<td>0.34</td>
<td>0.43</td>
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<td>0.83</td>
<td>0.69</td>
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<td>0.06</td>
<td>0.57</td>
<td>0.56</td>
<td>0.16</td>
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<tr>
<td>S -2</td>
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<td>32.96</td>
<td>32.73</td>
<td>32.59</td>
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<td>65.98</td>
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<td>66.56</td>
<td>66.90</td>
<td>61.27</td>
<td>63.48</td>
<td>66.67</td>
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<tr>
<td>Total</td>
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<td>100.04</td>
<td>100.40</td>
<td>99.66</td>
<td>100.77</td>
<td>100.99</td>
<td>100.01</td>
<td>100.53</td>
<td>100.04</td>
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<tr>
<td>Fe +2</td>
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<td>0.01</td>
<td>0.01</td>
<td>0.03</td>
<td>0.01</td>
<td>0.01</td>
<td>0.08</td>
<td>0.04</td>
<td>0.01</td>
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<tr>
<td>Cd +2</td>
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<td>0.00</td>
<td>0.00</td>
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</tr>
<tr>
<td>Mn +2</td>
<td>0.00</td>
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<td>0.01</td>
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<td>1.97</td>
<td>1.99</td>
<td>1.98</td>
<td>2.00</td>
<td>1.97</td>
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</tbody>
</table>

1. NKF-M-13-87, core of sphalerite crystal occurring in sulfide stringer in fine-grained admixture of rhodocrosite, rhodonite, and quartz.
2. NKF-AA-27-87, core of sphalerite crystal occurring in sulfide stringers in fine-grained admixture of rhodocrosite, rhodonite, and quartz.
3. NKF-BM-65-87, sphalerite crystal occurring in fine-grained gold-bearing sulfide seam containing pyrite.
4. NKF-BM-67-87, sphalerite crystal occurring in fine-grained gold-bearing sulfide seam containing pyrite + magnetite + hematite.
5. NKF-X-24-87, coarse-grained lemon-yellow sphalerite crystal of late base-metal mineralization.
6. NKF-R-18-87, colorless sphalerite in coarse-grained banded crystal occurring in veinlet cutting earlier gold-bearing sulfide seam.
7. NKF-R-18-87, red-brown sphalerite in coarse-grained banded crystal occurring in veinlet cutting earlier gold-bearing sulfide seam.
8. NKF-R-18-87, sphalerite from earlier gold-bearing seam containing magnetite + hematite + pyrite, cut by veinlet containing coarser sphalerite (nos. 6 and 7).
9. GSP-MN-87, coarse crystals of late-stage yellow sphalerite rimmed with minor chalcopyrite.

GYPSUM

Large curls of gypsum (CaSO₄·2H₂O) ranging in size from a few to tens of centimeters are found in open vugs and cavities along the vein structures. Gypsum having a fine-grained spongelike texture also occurs in vugs and cavities. These crystals certainly formed later than all of the ore-stage minerals and probably are of recent (modern) age.

HEMATITE

Hematite (Fe₂O₃), the most abundant iron oxide mineral, occurs in both mineral associations of the North Amethyst paragenesis. It is intergrown with precious- and base-metal sulfides and other iron oxides in the earlier association and with quartz, chlorite, and base-metal sulfides in the later association. Much of the hematite is coated with iron oxides; microprobe analyses of visually unoxidized grains indicate that they contain less than 1 weight percent combined MnO + SiO₂ + Al₂O₃ + MgO.

FIGURE 7.—Ternary diagram showing compositions of carbonates for three stages of the North Amethyst vein mineralization. Bars show range of compositions for carbonates present in more than one stage. Possible tie-lines are dashed.
Blades of hematite, 0.1 to 1.0 mm long, in electrum-bearing samples of the earlier association are intergrown with rounded anhedral magnetite grains (<50 μm in diameter) and cubes of pyrite. In some sections, the hematite is replaced by magnetite, while in other areas, the hematite has been altered to fine-grained iron oxides. Abundant hematite colors some ore samples a deep red hue.

Thin, transparent-red hematite flakes having well-developed hexagonal outlines occur in the later association as fine- to medium-grained crystals (<60 μm wide) intergrown with quartz, chlorite, pyrite, sphalerite, galena, and chalcopyrite. The flakes are also included in sphalerite and quartz crystals, where they frequently are clustered along colored growth zones.

**MAGNETITE**

Magnetite (FeFe₂O₄) is found in the earlier mineral association of the North Amethyst paragenetic sequence and is primarily associated with hematite and pyrite in a quartz-rich gangue. It occurs as equant grains (<0.5 mm wide) intergrown with euhedral pyrite and laths of hematite, and as a replacement of hematite. Some magnetite contains blebs of electrum several micrometers wide. Electron microprobe analyses of magnetite intergrown with hematite indicate virtually no substitution of MnO, Al₂O₃, SiO₂, ZnO, or MgO. Magnetite is not found with chlorite. In general, magnetite appears to be restricted to the earlier mineral association, whereas chlorite characterizes the later, base-metal association.

**MANGANESE SILICATE MINERALS**

At least four Mn-silicate minerals occur in the earliest stage of the North Amethyst ores. Rhodonite is the most abundant, and pyromangite, caryopilite(?), and inesite occur in much smaller quantities. The complete assemblages include rhodocrosite and quartz. Kutnohorite and Mn-calcite are also present in small amounts.

**Caryopilite** (Mn₆Si₄O₁₀(OH)₈).—Fine-grained intergrown aggregates of a pleochroic yellow to brown hydrated layer silicate (~<0.01×0.05 mm) are found in small, matted, 2-mm-wide clumps in the early, Mn-Au association. The mineral fills in around earlier formed quartz crystals and rims pyrite, chalcopyrite, and sphalerite stringers in a rhodonite + quartz admixture. The mineral clearly formed late, probably by retrograde alteration of rhodonite, pyromangite, and inesite. The mineral has been tentatively identified as caryopilite on the basis of petrographic characteristics as described in Abrecht (1989) and energy dispersive analysis.

**Inesite** ([Ca₄Mn₇Si₁₀O₄₆(OH)₂·5H₂O]).—Acicular crystals of inesite (table 7) up to 2 cm in length form radiating...
sheaths that fill fractures cutting silicified wall rock and early Mn-carbonate- and Mn-silicate-bearing assemblages in higher levels of the North Amethyst workings. Minerals intergrown with the coarse pale-pink to cream-colored crystals include quartz, rhodocrosite, Mn-calcite, galena, sphalerite, pyrite, and pyrargyrite.

**Pyroxmangite** \((\text{Ca}_{0.4-0.9} \text{Mn}_{0.86-0.92} \text{Fe}, \text{Mg}_{0.1-0.95} \text{Si}_3 \text{O}_8)\). — Pyroxmangite (table 7, fig. 10) is present in the earliest stage of mineralization. The mineral ranges in size from a few micrometers to, at most, 100 \(\mu\)m in length and is tentatively identified by X-ray diffraction analysis, petrographic study, and electron microprobe analysis but has not been confirmed by single crystal techniques. It is intergrown primarily with rhodocrosite, rhodonite, and quartz. Microprobe analyses of the mineral (table 7) show a small variation in calcium, iron, and manganese.

**Rhodonite** \((\text{Ca}_{0.8-0.9} \text{Fe}, \text{Mg}_{0.1-0.95} \text{Mn}_{0.86-0.92} \text{Si}_3 \text{O}_8)\). — Rhodonite is the most abundant Mn-silicate present in the early stages of the paragenesis (table 7, fig. 10). The mineral occurs primarily in a cryptocrystalline admixture of quartz, carbonate minerals, and other Mn-silicates and as coarser crystals edging small quartz-filled vugs. The coarser grained crystals range up to 50 \(\mu\)m in width and 200 \(\mu\)m in length.

**Marcasite**

Characteristic reflection pleochroism and strong anisotropy distinguish marcasite (FeS\(_2\)) from other iron sulfides. The mineral occurs primarily as 50- to 100-\(\mu\)m-wide tabular or carrot-shaped crystals growing on pyrite and, at times, is coated with chalcopyrite. Anhedral marcasite is found coating and filling cracks in pitted crystals of pyrite. Marcasite generally appears to have formed late in the base-metal–silica association, as carrot-shaped crystals resting on pyrite, as anhedral fillings, and as small fragments caught up in a fine quartz sediment matrix. Distribution of marcasite within the veins is difficult to define because of the fine grain size and scarcity of this mineral.

**Polybasite**

Polybasite \(((\text{Ag}, \text{Cu})_{16} \text{Sb}_2 \text{S}_n)\) is most commonly associated with acanthite and generally formed late with respect to other Ag-bearing minerals. The sulfosalts is admixed with galena and sphalerite and intergrown with chalcopyrite; it also replaces sphalerite. The softness of the mineral, light etching, and darker color adjacent to galena help to distinguish it from pyrargyrite.

**Potassium Feldspar**

Feldspar \((\text{K}_{0.96-0.92} \text{Na}_{0.02-0.08} \text{AlSi}_5 \text{O}_{18})\) occurs as anhedral to subhedral grains extensively replaced by fine-grained Mn-carbonate, rhodonite, and other Mn-silicates and by quartz in the earliest Mn-rich vein stage. The feldspars vary in composition from Or\(_{93}\) to Or\(_{96}\) and have barium contents that vary from ~0.1 to 2.4 weight percent BaO. These feldspars are compositionally similar to wall-rock phenocrysts of sanidine or orthoclase adjacent to the veins but locally have higher barium contents. The feldspars may have been incorporated into the vein during formation of the earliest stages of mineralization by metasomatic alteration of the volcanic wall rock. Potassium feldspar also occurs as 0.1- to 1.0-mm-long euhedral barium-rich crystals of adularia intergrown with amethystine quartz and filling small vugs, cavities, and fractures in most vein material. The adularia is associated with minerals of the base-metal association and is relatively common throughout the ore zone. Adularia ranges in content from Or\(_{92}\) to Or\(_{98}\) and contains 0.4 to 1.6 weight percent BaO.

**Pyrargyrite–Proustite**

Pyrargyrite \((\text{Ag}_3 \text{SbS}_3)\)-proustite \((\text{Ag}_3 \text{AsS}_3)\) primarily occurs alone in small (>10 \(\mu\))m anhedral grains scattered through a silicious gangue and in aggregates with acanthite that border galena and sphalerite. It also is intergrown with acanthite and native silver in aggregates that border or are enclosed by chalcopyrite. Other grains are
VEIN MINERALOGY AND COMPOSITIONAL RANGES

intergrown with acanthite and uyttenbogaardtite and rim chalcopyrite, tetrahedrite, and galena. Subhedral prismatic to anhedral crystal aggregates are enclosed by rhodochrosite and quartz and occur along cracks in broken pyrite cubes.

PYRITE

Pyrite (FeS₂) is the most abundant and ubiquitous sulfide mineral, as it occurs in almost all stages of both mineral associations. It is primarily associated with either quartz-chlorite-hematite assemblages or other base-metal sulfides. The pyrite shows idiomorphic habit, and cubes are the most common shape, although some overgrowths and aggregates are anhedral. Thin botryoidal crusts of pyrite occur late in the paragenetic sequence in places.

In the early association, subhedral equant pyrite crystals are bordered by sphalerite, rimmed by chalcopyrite and galena, and contain inclusions of all three sulfides. The crystals range in size up to a millimeter on edge; however, most are in the range 0.1 to 0.3 mm. Some of these crystals also contain rounded blebs of electrum. Grain margins of the pyrite crystals are often corroded and replaced by sphalerite and galena.

Some crystals are isotropic to faintly anisotropic and twinned. There is often a zonal arrangement with sphalerite, acanthite, and pyrargyrite crystals, which suggests that pyrite began to crystallize early, and its deposition continued as other sulfides and sulfosalts began to form. Later pyrite overgrowths on these crystals do not show inclusions. Rarely, marcasite crystals sit on pyrite. Pyrite also occurs as isolated equant cubes in quartz and carbonate gangue.

In the later association, euhedral pyrite is found in mixtures of quartz, hematite, and chlorite and is also intergrown with euhedral sphalerite and anhedral galena and chalcopyrite. The pyrite crystals of the late association are much larger in size, and crystals up to a centimeter on edge do occur. However, most pyrite crystals in this association are a millimeter or less on a side. Pyrite crystals in the size range 0.1 to 0.5 mm dust singly terminated quartz crystals and coarse crystalline calcite (fig. 5E) in the latest stage. They also occur as millimeter-thick layers within crusts of carbonate minerals. At times, 50- to 100-µm-sized carrot-shaped marcasite crystals overlie this latest stage pyrite.

SILVER

Native silver (Ag⁰) occurs primarily as scarce, lone 10- to 100-µm-sized grains in fractures and as millimeter-long wire aggregates in cavities or resting on broken surfaces. It is also found as scarce grains bordering galena that are enclosed by gangue and are rarely bordered by acanthite. In some samples, native silver borders sphalerite in aggregates that include chalcopyrite, jalpaite, and possibly stromeyerite. Textural relations indicate that native silver occurs much later in the paragenetic sequence than most other silver minerals.

SPHALERITE

Sphalerite ((Zn,Fe,Mn,Cd)S) is one of the most abundant sulfide minerals in the deposit. The mineral is found in two stages of the early, Mn-Au association and in one stage of the later, base metal-silica association. Sphalerites in different stages vary in grain size, color, and composition.
Stage alpha.—Small (<2 mm) rounded grains of intensely chalcopyritized sphalerite occur in small stringers within the cryptocrystalline admixture of Mn-silicates, carbonate minerals, and quartz that make up the earliest stage. The sphalerite is intergrown with cubes of pyrite and anhedral galena and contains generally low and subequal amounts of iron, cadmium, and manganese (table 8, Nos. 1 and 2).

Stage beta.—Abundant sphalerite is present also in the precious- and base-metal seams of the Mn-Au association. It is often rimmed and replaced by electrum, pyrrargyrite, acanthite, chalcopyrite, uytenbogaardtite, and galena. Sphalerite corrodes and sometimes replaces pyrite. Small equant grains also occur in siliceous gangue, with sparse micrometer-sized exsolution blebs of chalcopyrite. Sphalerite is bordered and corroded by all of the other sulfides, especially galena and chalcopyrite, and by tetrahedrite. All sphalerite of the Au-bearing stage is extensively chalcopyritized, and most crystals (table 8, No. 3) have a composition similar to that of the sphalerite described above. An important exception to this is that sphalerite intergrown with magnetite and hematite has a more iron rich composition (table 8, Nos. 4 and 8), generally containing around 2 percent iron by weight.

Stage 1.—Sphalerite crystals of the later, base-metal stage range in width from about a millimeter to a maximum of about 2 cm and range in hue from colorless to greenish to pale yellow to dark red-brown. The subhedral, equant crystals form aggregates that are intergrown with galena, euhedral pyrite, chalcopyrite, and minor tetrahedrite. All sphalerite of this association is chalcopyritized; it contains numerous microscopic inclusions of chalcopyrite that are especially abundant at the rims. Some grains also contain abundant minute blebs of tetrahedrite that are concentrated at grain boundaries.

While most grains are uniform in color, many larger crystals of sphalerite are color banded on a simple scale. The colors fall into three groups—red-brown, yellow, and colorless—that show only minor color variation. On the scale of individual crystals, the contact between colors is sharp rather than gradational.

The majority of the banded samples studied have colorless or yellow cores and red-brown rims (thin-section thicknesses ranging from 60 μm to 1 mm). Rims, in shades ranging from moderate red-brown to deep red-brown, are more finely banded than cores; however, they still do not show the beautifully well-developed banding of samples typical of the OH vein in the main Creede district as described by Barton and others (1977) and Roedder (1977). Other samples show an alternating set of color bands, from red-brown to yellow to red-brown, but these are less common. No attempt was made to correlate color banding in the sphalerite, because possible marker horizons lacked continuity.

Analyzed sphalerite from the later, sulfide-rich stage displays higher and more variable iron contents (up to 8 mole percent FeS) than earlier sphalerites (table 8, Nos. 5–8). Some of the darkest sphalerites were not analyzed, so the true range may be higher. Measured iron contents are not as high as those of sphalerite from the OH vein of the main mining district (Barton and others, 1977), which contains up to 16 mole percent FeS. However, the most typical iron contents of North Amethyst sphalerite (~1 mole percent FeS) are similar to the most typical iron contents of OH sphalerite (~1 to 2 mole percent FeS).

STEPPANITE
A silver sulfosalt resembling stephanite (Ag₅SbS₄) is intergrown with pyrrargyrite and polybasite in aggregates in a siliceous matrix. It is also found alone filling cavities and fractures in the siliceous matrix.

TETRAHEDRITE-TENNANTITE
Tetrahedrite-tennantite ([(Cu,Fe,Ag,Zn)₁₂(Sb,As)₄S₁₃]) occurs in all sulfide-bearing stages, usually in association with sphalerite, galena, pyrite, and chalcopyrite. Reported to be Ag-rich (S.W. Caddey, C.B. Byington, and D.M. Vardiman, written commun., 1988), energy dispersive analysis confirms that much of the tetrahedrite contains appreciable amounts of silver and arsenic.

In the early association, tetrahedrite is most often adjacent to sphalerite and galena and also is in aggregates with chalcopyrite and acanthite. Scarce anhedral grains of tetrahedrite occur marginal to, and intergrown with, chalcopyrite and acanthite. Tetrahedrite also occurs with polybasite, intergrown with pyrrargyrite and acanthite, alone, and intergrown with galena and chalcopyrite in cavity fillings.

In the later association, 50- to 70-μm-sized grains of tetrahedrite most often are with contemporaneous chalcopyrite and galena in aggregates that border sphalerite. Rounded micrometer-sized grains of tetrahedrite occur in sphalerite in a texture similar to that of chalcopyrite disease.

Growth banding was not observed in any of the samples, possibly because of their small grain size. Growth banding in tetrahedrite from the Bulldog vein of the Creede district is described by Plumlee (1989). He found that the growth bands reflected compositional variations in Ag, As, and Sb.

UYTENBOGAARDTITE
Uytenbogaardtite (Ag₃AuS₄) most often is intergrown with acanthite containing micrometer-sized electrum
than 2 mole percent combined FeCO$_3$ + MgCO$_3$. This in excess of 500°C, for rhodocrosite intergrown with and Peters (1981) at 540°C and a composition of Cc$_{32}$ Rc$_{68}$.

kutnahorite in the early assemblage (fig. 8). The most blage and temperatures in excess of 425°C, and possibly the calcite-kutnahorite compositions of the early assemblage. Upon cooling, the argentite would exsolve}

Ag-sulfide equilibria for the North Amethyst ores most likely reflect exsolution during cooling of a higher temperature Au-bearing argentite + Ag-rich electrum assemblage. Upon cooling, the argentite would exsolve uytenbogaardtite and leave the original Ag-rich electrum with acanthite and uytenbogaardtite. Thus, the Au- and Ag-sulfide equilibria for the North Amethyst ores most likely reflect subsolidus reequilibration at quite low temperatures and, like the Cu-Ag-sulfides, cannot be used to yield reliable temperatures of specific conditions of ore deposition because of their fast reaction rates.

DISCUSSION OF MINERAL EQUILIBRIA

GOLD AND SILVER SULFIDES

Experimental work by Barton (1978) demonstrated that uytenbogaardtite is stable at relatively low temperatures (<200°C) with Au-bearing argentite but that only below 113°C can uytenbogaardtite coexist with acanthite. Barton (1978) also showed that uytenbogaardtite is stable with relatively Au-rich electrum compositions (0.8-0.9 atomic fraction Au). The intergrowth textures described in this study for uytenbogaardtite, acanthite, and electrum are similar to those described by Barton and others (1978) for uytenbogaardtite from the Comstock lode. For that deposit, they suggested that the textures may reflect exsolution during cooling of a higher temperature Au-bearing argentite + Ag-rich electrum assemblage. Upon cooling, the argentite would exsolve uytenbogaardtite and leave the original Ag-rich electrum with acanthite and uytenbogaardtite. Thus, the Au- and Ag-sulfide equilibria for the North Amethyst ores most likely reflect subsolidus reequilibration at quite low temperatures and, like the Cu-Ag-sulfides, cannot be used to yield reliable temperatures of specific conditions of ore deposition because of their fast reaction rates.

CARBONATE MINERALS

Peacor and others (1978, 1987) have suggested possible solvii between calcite and kutnahorite and kutnahorite and rhodocrosite to account for the compositions of some naturally occurring carbonate minerals. Comparison of those solvii with the North Amethyst carbonate data suggests temperatures of formation above 350°C for the calcite-kutnahorite compositions of the early assemblage and temperatures in excess of 425°C, and possibly in excess of 500°C, for rhodocrosite intergrown with kutnahorite in the early assemblage (fig. 8). The most Ca-rich rhodocrosite of the earliest carbonate-bearing stage has a composition of Cc$_{23}$ Rc$_{75}$ and contains less than 2 mole percent combined FeCO$_3$ + MgCO$_3$. This composition is somewhat less calcic than the critical point of the CaCO$_3$-MnCO$_3$ solvus as reported by deCapitani and Peters (1981) at 540°C and a composition of Cc$_{38}$ Rc$_{62}$. (Experiments were run at 2 and 10 kbar, and no significant differences due to pressure were detected.) These temperatures are exceedingly high and, as discussed below, unreasonable for this environment.

Temperatures above 300°C are unusual for mineralization occurring in the typically shallow epithermal environment (Heald and others, 1987) where depths are generally on the order of 500 m. However, for the North Amethyst vein, about 1,500 m of volcanic and volcanioclastic rock may have overlain the base of Au-rich mineralized rock (Lipman and Sawyer, 1988; Foley, 1990). A simple analysis based on temperature-depth calculations of Haas (1971) indicates that the boiling point of a nonsaline fluid at 1,500 m is about 320°C; fluids that are more saline would boil at higher temperatures. Temperatures as high as those suggested by carbonate mineral equilibria are unlikely, however, because they would require formation at unreasonably great depths. The excessively high temperatures suggest that compositions of rapidly deposited carbonates of the early stage may be metastable.

Rhodocrosites and calcites from all later stages of mineralization have more restricted compositional ranges and lie outside experimental solvii (Goldsmith and Graf, 1957; dicapitani and Peters, 1981) at temperatures of less than 400°C.

MANGANESE SILICATE EQUILIBRIA

Pyroxmangite and rhodonite are polymorphs of MnSiO$_3$. Rhodonite is the high-temperature form and is stable above about 350°C at pressures of 1 to 2 kbar, whereas pyroxmangite of MnSiO$_3$ composition is the stable phase at low pressures below 350°C (Maresch and Mottana, 1976). The presence of both rhodonite and pyroxmangite can be used to constrain temperature if the compositions do not depart markedly from MnSiO$_3$ and if pressure is known. In this case, because up to 11 weight percent CaO and 2 weight percent FeO + MgO replaces MnO, and because pressures are not well constrained (Foley, 1990), the assemblage can be used only qualitatively. To a first approximation based on ionic radii, the substitution of Mg$^+$ and Fe$^+$ would stabilize pyroxmangite at higher temperatures, whereas the substitution of Ca$^+$ should stabilize rhodonite to lower temperatures (Maresch and Mottana, 1976).

If an ideal ionic activity model is used as a first approximation to describe the activity of MnSiO$_3$ in rhodonite and pyroxmangite and the phase equilibria data of Maresch and Mottana (1976) are used, the free energy of reaction can be written as:

$$
\Delta G_r = \Delta H_r - T \Delta S_r + P \Delta V_r + R T \ln K_r = 1347(\pm 100) - 2.069(\pm 0.2)T(\text{K}) + 0.0418(\pm 0.025)P(\text{bars}) + 8.3143 T(\text{K}) \ln \left( \frac{X_{\text{Mn, rhodonite}}}{X_{\text{Mn, pyroxmangite}}} \right)
$$
where $\Delta G_r = \text{Gibbs free energy of formation for the reaction}$

$\Delta H_r = \text{enthalpy of formation of the reaction}$

$\Delta S_r = \text{entropy of formation of the reaction}$

$P = \text{pressure}$

$\Delta V_r = \text{change in volume of the reaction}$

$R = \text{gas constant}$

$T = \text{temperature of the reaction}$

$K_r = \text{equilibrium constant for the reaction}$

$X = \text{mole fraction}$

This can be further recast to define temperature:

$$T(\text{°C}) = \frac{162(\pm 12) - 5.027(\pm 3.0)P(\text{kbars})}{0.2488(\pm 0.025) - \ln(X_{\text{rhodonite}}/X_{\text{pyroxmangite}})} - 273$$

When data for coexisting rhodonite-pyroxmangite pairs ($P = 0.5$ kbar) are used, an approximation to the temperature of equilibrium can be calculated:

<table>
<thead>
<tr>
<th>$T(\text{°C})$</th>
<th>$X_{\text{MnSiO}_3}$</th>
<th>$X_{\text{MnSiO}_3}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$327 \pm 58$</td>
<td>0.89±0.01</td>
<td>0.875±0.01</td>
</tr>
<tr>
<td>$339 \pm 58$</td>
<td>0.86</td>
<td>0.85</td>
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</table>

Temperatures at the middle to lower range of this approximation to the temperature of equilibrium (~330±60°C) are in general agreement with mineral assemblages, fluid-inclusion data, and other geologic considerations for the Mn-Au association of the North Amethyst vein system.

**MINERAL ASSOCIATIONS AND ASSEMBLAGES**

The ore and gangue minerals of the North Amethyst vein system are paragenetically complex; they fall into two mineralogically, texturally, and chemically distinct associations. The associations overlap spatially but were separated in time by a period of extensive brecciation and sedimentation. The earlier association is characterized by two mineralogic stages, whereas the second, later association is subdivided into three stages on the basis of mineralogy and textural relations. Successive stages in each association are generally marked by relatively sharp rather than gradational changes in mineralogy and by periods of movement, brecciation, sedimentation, or replacement. In only a few instances are the transitions characterized by intergrowths or interbanding between mineral assemblages of adjacent stages.

Figure 11 describes the sequence of events relating to formation of mineralized rock in the North Amethyst area. The general sequence of mineral deposition is shown in figure 12. Temporal and spatial relationships between mineralization, brecciation, and sedimentation are shown in a series of diagrams beginning with figure 13.

**ALTERATION AND PRE-ORE VEIN ASSEMBLAGES**

Several types of wall-rock alteration have been observed in the North Amethyst vein system. These include intense sericitic alteration at higher levels in the system, two stages of potassium metasomatism of the wall rock at depth, and a bleaching of the wall rock adjacent to some vein structures at all levels (figs. 11, 13). A detailed examination of the alteration assemblages was beyond the scope of this study and is unwarranted because of existing regional studies of alteration relations in the Creede area. However, certain observations can be made on the basis of field relations, hand-sample study, geochemical variations, and some X-ray diffraction analyses.

**SERICITIC ALTERATION**

Intense sericitic alteration characterized by illite-smectite clay minerals is generally restricted to above the 10400 level of the North Amethyst vein workings (see fig. 6). Geochemical analysis of selected material by the WDXRF and graphite AA techniques outlined in Baedecker (1987) (N.K. Foley, unpub. data, 1990) shows that the most intense sericitically altered rock consists of about 90 percent SiO$_2$ + Al$_2$O$_3$, 4 percent K$_2$O, and 2.5 percent combined Fe$_2$O$_3$ + FeO + MgO; measured precious-metal values (~5 ppm Au) are much higher than those of typical unaltered volcanic rock but lower than those of substantially less-altered wall rock adjacent to veins lower in the workings (20 to 40 ppm Au).$^5$

The light-colored sericitic alteration is cut by seams of fine-grained dark-colored clay (mixed illite-smectite by X-ray diffraction). Quartz veins also cut the sericitic material; sharp contacts suggest that the veins formed when the rock was more competent, prior to the peak alteration event. Clear-cut paragenetic relations between ore mineralization and this intense alteration were not found; however, the alteration zone appears to overlie the zone of base-metal-rich mineralized rock and cuts across the zone of Au-bearing mineralized rock. Au- and Mn-silicate-bearing assemblages occur at elevations above and below the intense alteration zone, but the base-metal-rich later stages are only below, as is true for veins in the main Creede district. No information is available on the presence or absence of alteration assem-

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$^5$ Average gold values are notoriously difficult to determine because of the effect of a single nugget of electrum on a bulk analysis, so these values are given only for general discussion. They do not reflect actual modes or means.
<table>
<thead>
<tr>
<th>Process</th>
<th>Early Association</th>
<th>(Time →)</th>
<th>Late Association</th>
</tr>
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<tbody>
<tr>
<td>Mineralization</td>
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<tr>
<td>K-metasomatism</td>
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<td>Bleach</td>
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<td>Early quartz/carbonate veins</td>
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<td>Stage alpha</td>
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<td>Stage beta-1</td>
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<td>Stage 1</td>
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<td>Fracturing and vein movement</td>
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<td>Brecciation</td>
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<td>Breccia-1 -2 -3 -4</td>
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<td>Sedimentation</td>
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<td>Leaching and replacement</td>
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</table>

**Figure 11.** — Progression of events in the development of the North Amethyst ores. See text for description of stages and named breccias. Arrow locates interval containing dated adularia.

Blazes above the highest elevations of gold mineralization. Rocks cropping out on the hillside above the North Amethyst portal are stained by iron oxides all along the trace of the Equity fault.

Well-developed clay alteration zones cap vein structures throughout much of the central and southern Creede district (Steven and Eaton, 1975; Barton and others, 1977; Horton, 1983). Barton and others (1977) and Bethke and Rye (1979) proposed that the caps formed primarily as a result of reaction of the wall rock with acidic fluid that resulted from condensation of ore fluid that boiled at depth. An age date on illite-smectite from the cap material in the OH vein of the southern part of the district is in agreement with dates measured on vein adularia and sericite (table 1). A similar interpretation could explain the relation between overlying alteration-zone and base-metal mineralization in the North Amethyst area; however, any relation to gold mineralization is left unexplained.

**POTASSIUM METASOMATISM AND BLEACHING**

Wall rocks of the North Amethyst vein are potassium metasomatized at depth. A general increase in potassium content of the wall rock with depth is reported by Sawyer and others (1989), who suggest that the intensity of potassium metasomatism may increase with depth in the vein system. This pattern may be influenced by type and distribution of veins within the wall rock, because bleached wall rock has elevated potassium contents over adjacent unbleached wall rock at all depths (N.K. Foley, unpub. data, 1990). Moreover, the barium contents of potassium feldspar in the bleached wall rock are higher than barium contents of unbleached wall rock, a fact suggesting that the bleaching event was chemically distinct from the regional pre-ore potassium metasomatic event (N.K. Foley, unpub. data, 1990). Sharp contacts between bleached and unbleached wall rock, coupled with the chemical distinctions cited above, suggest that the bleaching event postdated the potassium metasomatism; however, this needs to be substantiated by studies that delineate the extent of the metasomatism within the Nelson Mountain Tuff and that determine mass transfer among the alteration zones and vein mineral assemblages. Within the central and southern Creede district, a bleaching event confined to wall rock adjacent to a vein does postdate the regional potassium metasomatism of the Carpenter Ridge Tuff (P. Barton and P. Bethke, oral commun., 1989).
The bleaching event (fig. 13A) resulted in a color change from gray and gray-tan to off-white of widths ranging from a few centimeters to tens of centimeters in wall rock adjacent to some veins. The event apparently predates all mineralization because vein minerals of both associations fill structures that cut both bleached and unbleached wall rock. In drillcore away from the underground workings, similar bleached fractures are filled with a fine-grained mixture of pyrite, quartz, and, at times, hematite and (or) chlorite.

Unbleached Nelson Mountain Tuff forms the wall for many of the mineralized veins. In general, the rock is hard and brittle, medium to dark gray, and retains volcanic textures consisting of phenocrysts (<2 mm) and flattened pumice in a very fine groundmass. The brittle nature indicates that the Nelson Mountain Tuff is silicified adjacent to veins, although chemical analyses of the rock show that SiO₂ and total alkali contents are not much elevated over values typical for volcanic rocks of dacitic-rhyolitic composition, on the basis of the IUGS classification scheme (LeMaitre, 1989; N.K. Foley, unpub. data, 1990). In the northern part of the workings, the veins cut slightly coarser grained rocks, possibly intrusive dacitic Nelson Mountain Tuff (Equity Member; Lipman and Sawyer, 1988), that are propylitized; they contain epidote, calcite, and abundant chlorite.

Most of the wall rock visible underground is cut by small (<3 cm) fractures that are crosscut in turn by minerals of the two main sulfide-bearing associations—the Mn-Au and base-metal-silica. These fractures are filled primarily with quartz and calcite and cut both bleached and unbleached wall rock. In drillcore away from the intensely mineralized area, bleached fractures contain mainly quartz, calcite, and pyrite, and occasionally barite and minor amounts of base-metal sulfides. In drillcore south of the exploratory workings, barite has

![Figure 12](image_url)
also been identified in similar fractures (D. Sweetkind, written commun., 1990). Barite was not found in samples collected underground and examined in this study, although it has been reported (S.W. Caddey, C.B. Byington, D.M. Vardiman, oral. commun., 1987). As described earlier, possible barite molds have been identified in samples of early assemblages collected from the underground workings.

MANGANESE-GOLD ASSOCIATION

The Mn-Au association is so named because of the presence of electrum and Au-Ag sulfide in one stage and because of the abundance of Mn-bearing minerals in the other stage. The occurrence and proportion of these two elements and the textural character of the minerals distinguish this association from the later, base-metal-rich association of the North Amethyst vein system and from other mineralized veins found throughout the Creede district.

The first stage, alpha, is characterized by Mn-carbonate minerals, Mn-silicates, and minor base-metal sulfides. The second stage, beta, has abundant precious- and base-metal minerals and minor quartz and carbonate gangue. Each stage has two mineralologically similar
substages identified by crosscutting relations. The sulfide-rich substages were separated by a period of fracturing and the development of a breccia having characteristic fragments.

STAGE ALPHA

The first vein-filling event that is directly related to ore development in the North Amethyst area occurred during faulting and fracturing of the volcanic wall rock. This event introduced Mn-rich fluids that deposited a fine-grained, almost cryptocrystalline, banded mineral mixture consisting predominantly of rhodocrosite, manganan calcite, rhodonite, pyroxmangite, and quartz (figs. 11, 14A). Locally, this fine-grained material of stage alpha coated a platy, tabular mineral (barite?) that was subsequently leached; the molds are filled with coarser adularia and quartz (probably stage-2 minerals; see below). Minor sulfides, primarily pyrite, sphalerite, and galena, occur in stringers in the Mn-carbonate, Mn-silicate, and quartz matrix. Retrograde alteration of this material produced a complex suite of hydrated Mn-silicate and Mn-carbonate minerals.

Vein widths of stage alpha vary from a few centimeters to more than a meter. Features that characterize this stage are best preserved in the narrower veins; wider veins show multiple fracturing events and recrystallization of minerals.

Alpha-1 substage veins cut bleached and unbleached wall rock and precursor veins, indicating that both the bleaching event and the development of precursor veins predated this stage. In general, contacts of vein and wall rock are sharp; however, the occurrence of embayed and partially replaced grains of potassium feldspar in the Mn-rich matrix suggests that some metasomatic replacement of wall rock may have occurred.

Deposition appears to have progressed from the walls toward the center of the veins, leaving open spaces in the center. The walls of the spaces have a rounded, bulbous texture and were filled later with 1- to 2-mm-long adularia rhombs, coarse amethystine quartz (~2 to 3 cm long), rhodocrosite (1 to 5 mm wide), and, in places, fluorite (1 to 2 mm wide) and sphalerite (1 to 2 mm wide; probably stage-2 minerals).

The alpha-2 substage consists of a fine-grained mixture of quartz, rhodocrosite, and Mn-silicate minerals deposited around fragments of alpha-1 substage and wall rock (fig. 15). This substage is more quartz rich than the earlier alpha substage and has a greater proportion of base-metal sulfides. Definition of this stage and its exact position in the paragenesis is complicated by evidence that most fluids of later stages reacted with or recrystallized stage alpha to some extent. Alpha-2 substage is probably a product of the local reaction of later stage fluids with fragments of the alpha-1 substage; however, a second influx of Mn-rich fluids may have contributed to the mineralization.

STAGE BETA

Continued movement of the North Amethyst vein and fracturing of the stage-alpha assemblage caused the vein system to open up again. Precious- and base-metal-rich fluids invaded the fractures and precipitated a complex suite of Au-, Ag-, Cu-, Zn-, Pb-, and Fe-bearing sulfides and sulfosalts (figs. 12, 14B). The location of the sulfide-rich seams may have been controlled in part by the structurally weak vuggy zones in the core of stage-alpha vein material. Seam widths vary from a few millimeters to tens of centimeters, although most samples studied had widths of 2 to 3 cm. The ore seams are reasonably intact in some areas, but most show minor offsets. In other areas, the seams are stretched and broken, and sulfide-rich pods are more strongly deformed.

Stage-beta fluids appear to have irregularly altered stage-alpha minerals from a dominantly Mn-carbonate, Mn-silicate, quartz assemblage to a Mn-carbonate and quartz assemblage. During the retrograde alteration, hanging-wall sides of the sulfide-rich seams are often recrystallized to a coarse Mn-calcite, whereas footwalls are altered to a dusty fine-grained mixture of rhodocrosite and quartz (fig. 4D).

The complex suite of precious- and base-metal sulfides precipitated by stage-beta fluids is listed in table 3, and the paragenetic sequence for the most common stage-beta minerals is shown in figure 12. The intergrowth of gangue, sulfides, and sulfosalts shows that, in general, deposition within stage beta was initiated by precipitation of pyrite and sphalerite, followed by galena, chalcopyrite, electrum, and other Au and Ag minerals. Although the depositional sequence was interrupted by minor periods of dissolution (for example, replacement of pyrite crystals; chalcopyrite, and tetrahedrite disease in sphalerite; corroded sphalerite crystals), stage-beta precipitation was generally continuous. Native silver, stromeyerite, and other minerals are found mainly in tiny late-stage veinlets that cut the stage-beta seams or as rims on earlier phases, and may be supergene.

Stage-beta seams show a mineralogic zonation that is discontinuous along strike of the seams and irregularly distributed across the vein widths. Zn, Fe, and Pb minerals occur across the width of all seams, while chalcopyrite and other Cu-bearing minerals are concentrated toward the center of the seams. In general, electrum and Au- and Ag-sulfides and sulfosalts occur with, or are generally formed later than, chalcopyrite.

The iron-oxide and -sulfide assemblage of stage beta consists of magnetite, hematite, and pyrite. This relatively oxidizing assemblage occurs locally along the
stage-beta seams and is most frequently found at depth in the system. The irregular distribution of the iron-oxide minerals, in contrast to the pervasive occurrence of pyrite, may indicate local change from pyrite equilibria to pyrite-hematite equilibria or a pyrite-magnetite-hematite buffer by an oxidative process such as boiling (Foley, 1990).

The pyrite, hematite and magnetite are intergrown with fine-grained Zn-, Pb-, and Cu-sulfides and abundant quartz. Electrum is the only precious-metal mineral commonly found with the iron-rich assemblage; it is intergrown with magnetite and pyrite and compositionally is similar to electrum in other assemblages. Euhedral magnetite is intergrown with pyrite and hematite in...
ORES OF THE NORTH AMETHYST VEIN SYSTEM, COLORADO

Disseminated base-metal sulfides in stage alpha-2

Unbleached wall rock

Stage alpha-2

Bleached wall rock

Alpha-1

FIGURE 15.—Two periods of Mn-carbonate and Mn-silicate mineralization of stage alpha are shown by the occurrence of one in the other.

apparent equilibrium, and magnetite also replaces hematite blades; the contrasting textures support a change from pyrite to pyrite-hematite-magnetite and locally to pyrite-magnetite or magnetite equilibria, as noted previously. Hematite appears, in some cases, to be altered to limonite; this alteration probably is supergene in origin.

The occurrence of rounded cobbles of stage beta in breccias and sedimented zones cut by later competent seams of stage-beta sulfides was the basis for defining two substages (beta-1 and beta-2). The two have essentially identical mineralogies and differ only in their occurrence (fig. 16). Lead-isotopic compositions of galenas from the two substages are identical within analytical error (Foley, 1990). Electrum compositions and sphalerite compositions are also virtually identical for the two substages (tables 6 and 8).

Breccia-1.—Precious-metal minerals were deposited in two pulses separated by a period of brecciation and sedimentation along parts of the vein. The breccia that developed between the two beta substages is characterized by an unsorted mix of subangular to rounded pieces of fine-grained Au-bearing ore, angular fragments of bleached and unbleached wall rock, and fragments of stage alpha held together with a microcrystalline, dominantly quartz cement (fig. 16).

TRANSITIONAL BRECCIAS AND SEDIMENTS

The transition between deposition of the Mn- and Au-rich assemblages and assemblages of the base-metal–silica association is marked by extensive brecciation of, and sedimentation in, the veins. The movement along the veins may have been a continuation, on a much larger scale, of the fracturing that formed breccia-1. Most of the breccias and sediments formed during this period are lumped together under the category breccia-2, with the exception of a matrix-supported, quartz-cemented breccia referred to as the “black quartz breccia.” The different breccia types record a short history of the transition period from the initial dynamic hydrology that reworked fragments and deposited them as fragment-supported breccias in large shoots and sediment “dumps” to the more quiescent hydrologic setting that resulted in graded bedding and chemically cemented, matrix-supported breccias.

BRECCIA-2

Breccias that are lumped together in this group have fragments that range in width from a few millimeters to 15 cm and in shape from angular to rounded (fig. 17). The fragments consist of all earlier stages of mineralization and wall rock; most fragments were derived from points close to the site of deposition.

Large breccia zones, resembling “shoots” of sedimented material with local graded bedding and quartz cement, formed as the fluids dumped their sediment load. The unsorted material forming the breccia zones consists mainly of subangular to rounded, bleached and unbleached fragments of volcanic wall rock, rounded and altered pieces of stage alpha, rounded to subrounded cobbles up to 15 cm in width of sulfide-rich stage beta, and fragments of earlier breccias (fig. 17A,B). The pieces are held together with quartz-rich cement. Interstitial sediment between larger fragments has graded bedding, soft-sediment deformation, and other textures indicating that the material settled out of solution after the larger cobbles and blocks were dumped (fig. 17B).

In another area, silt- to gravel-sized fragments settled out in large graded beds of at least four cycles that extend for 30 m along a vein wall and are greater than 3 m in height (tops and bottoms were not visible; fig. 4G, 17C). The water-deposited beds show minor soft-sediment deformation and an occasional sulfide-bearing cobbles (up to 8 cm in width) that has fallen into finely graded sediment (fig. 4H). The graded beds are truncated at one end by a mineralized linking fault, and at the other end they appear to drape against hanging-wall volcanic rocks.

Black quartz breccia.—Late in this transition period, veins of “black quartz breccia” developed along planes of movement (fig. 17D). The black quartz breccias consist of 50 to 90 volume percent microcrystalline quartz-rich matrix and 10 to 50 percent fragments of stage-beta and stage-alpha material, earlier breccias, and wall rock. The
cement is composed dominantly of microcrystalline quartz having a granular texture with sutured grain boundaries. The cements grade in color from light gray to black, apparently in relation to the amount of fine sulfides and electrum contained therein. It has not been possible to ascertain whether the sulfide aggregates are clastic or if they formed as precipitates, but the shapes of the grains suggest that they were broken and ground. They may be finely ground dust from earlier stage-beta sulfide seams.

BASE-METAL–SILICA ASSOCIATION

This association has three mineralogical stages: (1) an interval consisting dominantly of base-metal sulfides, iron oxides, and quartz; (2) an interval of Ca-, Mn-, and Fe-bearing carbonate minerals, fluorite, and quartz; and (3) a Mn-calcite, quartz, and pyrite interval. The association is named after the most abundant constituents, base-metal sulfides and quartz. Complete mineralogical listings of all stages are given in table 3.
STAGE 1

The first stage of the base-metal-silica association (fig. 18) is characterized by abundant base-metal sulfides, argentian tetrahedrite, hematite, chlorite, and quartz. Two mineralogically distinct substages are recognized: a chlorite, hematite, and quartz assemblage was deposited prior to, and concurrently with, the major period of base-metal mineralization that resulted in a crustified sequence of sphalerite, galena, pyrite, chalcopyrite, and tetrahedrite. The fine-grained chlorite, hematite, and quartz assemblage was deposited on blocks of earlier breccias and in fractures cutting earlier assemblages of the Mn-Au association. Isolated crystals of sphalerite appear in the siliceous mixture. Some of the siliceous mixture was fragmented and recemented before the fluids began depositing relatively coarse base-metal sulfides. In other areas, the sulfides are intergrown with chlorite, hematite, and quartz without obvious breaks, so a gradual shift in dominant mineralogy is suggested.
The coarse-grained sphalerite, galena, pyrite, and chalcopyrite assemblage of stage 1 contains interstitial quartz, chlorite, and hematite. Hematite flakes also are included in the sphalerite crystals. No clear-cut paragenetic relations are apparent among the minerals, with the exception that fluid reacted with sphalerite along fractures and grain boundaries to form chalcopyrite- and tetrahedrite-disease textures (Barton and Bethke, 1987). Sphalerite crystals are corroded, and periods of dissolution and reprecipitation are identifiable through studies of thin sections. Late-forming galena of stage 1 occurs as skeletal crystals in vugs that were subsequently filled with minerals of stage 2 (fig. 18).

Breccia-3.—Continued local fracturing of stage-1 minerals formed a breccia that is characterized by fragments of coarse base-metal sulfide and chloritized wall rock cemented by a slurry of cryptocrystalline quartz, chlorite, and pyrite (figs. 5F, 19). The chlorite gives the...
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Stage beta-2

Stage-beta fragment

Breccia-3, base-metal sulfide fragments, quartz, and chlorite cement

Relatively unbroken Stage 1

Breccia-3

Breccia-1

Breccia-2

Stage beta-2

Figure 19.—Local development of breccia-3, characterized by fragments of stage-1 sulfides and a green, chlorite-rich cement.

Cement a green hue. Minor amounts of base-metal sulfides were deposited on this brecciated material. Fragments of the base-metal sulfide are angular to subangular, and the wall-rock fragments are rounded. The breccia is most frequently found where stage-1 vein widths are greatest.

Stage 2

The second stage of mineralization in the base-metal-silica association (fig. 20) is characterized by coarse-grained quartz, calcite, rhodocrosite, and fluorite, all intergrown with minor amounts of adularia, chlorite, and pyrite. Locally, manganosiderite occurs within the calcite. Stage 2 was deposited on breccia-3 and stage 1. The minerals are coarse grained; quartz crystals grew up to 40 mm in length, while green fluorite cubes attained 13 mm on a side. Smaller octahedra of fluorite (<3 mm) are colorless or pale green or purple. In general, quartz, chlorite, and calcite were deposited before fluorite and additional quartz. Quartz crystals are zoned from colorless and white to amethystine and contain solid inclusions of calcite, adularia, and fluorite. Occasionally, a small amount of fine gray quartz sediment was deposited in vugs on crystals of stage 2.

Quartz and adularia that fill barite(?) molds in stage alpha probably crystallized from stage-2 fluids. The minerals are texturally identical to stage-2 minerals and have fluid-inclusion salinities and homogenization temperatures that are similar to those of other stage-2 minerals (Foley, 1990). Rhodonite crystals (up to 0.5 mm in length) that form selvages and project into some of the quartz-filled molds, and are intergrown with some rhodocrosite, may have formed from recrystallization of finer grained stage-alpha material.

Breccia-4.—This breccia contains angular fragments of all earlier assemblages, some breccias, and broken crystals and is cemented with a pale- to light-pink calcium-rich carbonate having a composition of Ca<sub>90</sub>Rc<sub>5</sub> (fig. 21). It is especially prevalent in the north parts of the vein system. The fragile and vuggy nature of the breccia suggests a late formation, probably concurrent with the development of stage 3.

Stage 3

The last clearly defined mineralization in the paragenetic sequence was deposition of quartz or manganocalcite, with pyrite, along the North Amethyst structure (fig. 22). Movement along the structure created vugs, cavities, and brecciated zones that cut all of the earlier stages. This movement was accompanied by, or followed by, leaching of earlier vein material along the entire length of the North Amethyst workings.

In the northern part of the ore zone, extensive crusts of pink to pale-yellow Mn-bearing calcite, interlayered with thin layers of crustiform pyrite, were deposited in the vugs and on material leached earlier. The carbonate crusts ranged in total thickness from millimeters to a meter. The carbonate minerals were selectively leached and now contain large, open vuggy zones.

Leaching of earlier stages, especially the extensive breccias, appears to have been more extensive in the southern workings. Samples of breccia-2 exhibit gradations in the degree of leaching ranging from bleaching of fragments to alteration of phenocrysts to clay minerals to samples that consist entirely of a foamy siliceous material containing molds possibly of leached blocks of wall rock and rare specks of dark sulfides. Geochemical analysis of the most altered and leached material indicates that, of the major elements, only SiO<sub>2</sub>(~94 percent), and minor Al<sub>2</sub>O<sub>3</sub>(~3 percent) and K<sub>2</sub>O (~1 percent) remain. Relatively high values of base and precious metals also persist (~90 ppm Au, 30 ppm Ag). This leached material is overgrown by large terminated crystals of quartz in the southern part of the ore zone. The quartz crystals are included in stage 3 because they overgrow the leached base; however, their formation may predate the manganocalcite. Minute cubes of pyrite dust the quartz surfaces; timing of this pyrite relative to the crustiform occurrence within manganocalcite is not clear.

Correlation of Mineral Assemblages with Other Vein Systems

The relation of the North Amethyst ores to those of the main Creede district 5 km to the south is of great
importance in assessing the nature, extent, and timing of mineralization. The mineralogy, textures, and paragenetic sequences established in this study can be used to compare the North Amethyst veins to veins of the main mining district at Creede. Paragenetic relations for the North Amethyst vein system and some veins of the main mining district are summarized in table 9. Mineral stages defined for the North Amethyst vein system can be correlated, in part, with mineral stages defined in previous studies for the southern Amethyst vein (Guidice, 1981; Robinson, 1981; and Robinson and Norman, 1984), the Bulldog Mountain vein system (Plumlee, 1989), and the OH and P veins (Barton and others, 1977; Bethke and Rye, 1979; P.B. Barton, Jr., oral commun., 1989). Most
of the corresponding stages are mineralogically similar and occur throughout the district; however, there are some important exceptions.

At present, stages alpha and beta of the Mn-Au association have been found only in the North Amethyst area. The assemblages have not been described elsewhere within the boundaries of the Creede mining district. Mn-silicates and uytenbogaardtite are not found south of the North Amethyst workings, although rare occurrences of electrum have been reported for a couple of localities. An occurrence of dark-yellow gold with a high degree of fineness contained in a gangue of manganese oxides has been described by Emmons and Larsen (1913, 1923) from the Happy Thought and Amethyst mines of the Amethyst vein. The gold occurred in cracks and veinlets cutting older ore and was said to be a common supergene occurrence in near-surface workings. Gold also has been reported as inclusions in pyrite, galena, sphalerite, and other minerals (Emmons and Larsen, 1923); however, no additional details are available. P.M. Bethke and P.B. Barton, Jr. (oral commun., 1989), note the occurrence of abundant finely disseminated gold in early chalcopyrite in a single sample from the OH vein (their stage A or B, table 9). Galenas from North Amethyst gold-bearing stage beta are distinct in their lead-isotopic character from galenas of stage 1 of the North Amethyst veins and from most veins of the main Creede mining district (Foley, 1990; Foley and Ayuso, in press). The composition of OH and P vein tetrahedrite is variable but generally is silver rich (L.B. Wiggins and T.L. Woods, unpub. data, 1980), similar to that of the North Amethyst ore.

Stage 2 of the North Amethyst vein correlates minerallogically with stage C of the OH and P veins (table 9). Both stage 2 and stage C contain fluorite, quartz, and carbonate minerals. The North Amethyst stage-2 carbonate minerals consist of intergrown calcite and manganosiderite, whereas the OH and P veins contain only manganosiderite (Wetlauffer, 1977).

OH and P vein stage D ores are apparently absent in the North Amethyst vein system (table 9). Stage D consists largely of coarsely crystalline sphalerite, galena, quartz, chalcopyrite, and hematite. D-stage sphalerite is free of chalcopyrite disease and is color zoned from core to rim with yellow-white to red-brown to yellow-white zones having iron contents of less than 3 mole percent FeS (Barton and others, 1977).

Stage 3 of the North Amethyst vein system can be correlated with stage E of the OH and P veins (table 9; defined as stages A through E). Stage A of the northern OH and P veins consists primarily of massive chalcedonic quartz intergrown with rhombs of adularia. This stage is not found in the North Amethyst ores; however, in the interval of brecciation and sedimentation between the two sulfide-rich North Amethyst stages (stage beta and stage 1), quartz was deposited as chalcedonic cement in breccia. Although not defined as a vein-filling stage, the quartz cement may correlate with OH vein stage-A quartz, and this correlation would indicate a districtwide period of silica deposition.

Stage B of the OH and P veins is dominated by fine-grained base-metal sulfides—sphalerite, galena, chalcopyrite, and lesser amounts of Ag-rich tetrahedrite—contained in a chlorite, pyrite, hematite, and quartz gangue assemblage. This stage correlates directly with stage 1 of the North Amethyst vein (table 9). Sphalerites of the North Amethyst, OH, and P vein assemblages show extensive chalcopyrite- and minor tetrahedrite-disease textures. Zoned sphalerites from the OH stage B and North Amethyst stage 1 are compositionally similar; light zones have generally low iron contents (<2 mole percent), and dark bands are iron rich (up to 8 mole percent for North Amethyst stage 1, up to 16 mole percent for OH stage B). Galenas (stage 1B) from all three veins also have similar lead-isotopic compositions (Foley, 1990; Foley and Ayuso, in press). The composition of OH and P vein tetrahedrite is variable but generally is silver rich (L.B. Wiggins and T.L. Woods, unpub. data, 1980), similar to that of the North Amethyst ore.

Stage 2 of the North Amethyst vein correlates minerallogically with stage C of the OH and P veins (table 9). Both stage 2 and stage C contain fluorite, quartz, and carbonate minerals. The North Amethyst stage-2 carbonate minerals consist of intergrown calcite and manganosiderite, whereas the OH and P veins contain only manganosiderite (Wetlauffer, 1977).

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Stage 3 of the North Amethyst vein system can be correlated with stage E of the OH and P veins mainly on the basis of the occurrence of pyrite in all of these stages (table 9). Pyrite in North Amethyst stage 3 is volumetrically minor compared to stage E, and quartz and
FIGURE 22.—Stage 3 is marked by deposition of thick crusts of a coarse pink calcite, pyrite, and quartz. Inset shows calcite and pyrite occurrence.
ores of the north amethyst vein system, colorado

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Table 9.—Districtwide correlation of mineral assemblages and brecciation events

<table>
<thead>
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<tbody>
<tr>
<td>(3) Mn-Ce + Qz + Py</td>
<td>(E) Py + Stib</td>
<td>(V) Py + Stib + Pyr</td>
<td>Absent?</td>
<td>(2) Py + Qz.</td>
</tr>
<tr>
<td>Breccia-4</td>
<td>Absent?</td>
<td>Absent?</td>
<td>Absent?</td>
<td>Absent?</td>
</tr>
<tr>
<td>Absent</td>
<td>(D) Sp + Gn + Qz + Cpy + Hm</td>
<td>(IV) Sp + Gn + Cpy + Td + Pear + Ac</td>
<td>Absent?</td>
<td>Absent?</td>
</tr>
<tr>
<td>(2) Qz + Mn-Ce + Fl + Re + Mn-sid + Chl + Py + Ad</td>
<td>(C) Fl + Qz + Mn-sid + Sp + Gn</td>
<td>Absent?</td>
<td>Absent?</td>
<td>Included in (1)?</td>
</tr>
<tr>
<td>Breccia-3</td>
<td>Absent?</td>
<td>Absent?</td>
<td>Absent?</td>
<td>Absent?</td>
</tr>
<tr>
<td>(1) Sp + Gn + Py + Cpy + Qz + Hm + Chl + Td</td>
<td>(B) Qz + Sp + Chl + Gn + Hm + Td + Au</td>
<td>(II) Bar + Qz + Sp + Gt + Td</td>
<td>Absent?</td>
<td>Absent?</td>
</tr>
<tr>
<td>Breccia-2</td>
<td>(A) Qz + Ad</td>
<td>(I) Re + Qz + Sp + Gt + Ad</td>
<td>Absent?</td>
<td>Absent?</td>
</tr>
<tr>
<td>(Beta-2) Gn + Sp + Cpy + Py + Td + Ac + Uyten + Au</td>
<td>Absent?</td>
<td>Absent?</td>
<td>Absent?</td>
<td>Absent?</td>
</tr>
<tr>
<td>Breccia-1</td>
<td>Absent?</td>
<td>Absent?</td>
<td>Absent?</td>
<td>Absent?</td>
</tr>
<tr>
<td>(Beta-1) Gn + Sp + Cpy + Py + Td + Ac + Uyten + Au</td>
<td>Absent?</td>
<td>Absent?</td>
<td>Absent?</td>
<td>Absent?</td>
</tr>
<tr>
<td>(Alpha) Re + Qz + Mn-Ce + Rh + Px + Sp + Gn + Py + Ad</td>
<td>Absent?</td>
<td>Absent?</td>
<td>Absent?</td>
<td>Absent?</td>
</tr>
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manganoan calcite occur in relatively greater proportions.

The sequence of mineral stages defined for the Bulldog Mountain vein system has been correlated with mineral stages of the OH and P veins by Plumlee (1989). The comments and ideas presented in that study may hold true for comparing the North Amethyst and Bulldog Mountain paragenetic sequences because of the mineralogical similarities between the OH and North Amethyst assemblages (table 9). As in the case of the OH vein, chalcedony of Bulldog Mountain stage I may correspond to chalcedonic cements of North Amethyst breccia-2. The North Amethyst stage 2 is more like the sphalerite-and galena-rich, Ag-poor assemblages of the northern Bulldog Mountain stage II than the barite and Ag-rich assemblages of the southern half of the Bulldog system. However, in contrast to the North Amethyst stage 2, northern Bulldog Mountain stage II does contain some barite and does not have abundant chalcopyrite-hematite-quartz gangue. Compositions of carbonate minerals of North Amethyst stage 2 (table 5) are similar to those of Bulldog Mountain stage III (Wetlaufer, 1977).

Thus, Bulldog Mountain stage III may correlate with North Amethyst stage 2. Bulldog Mountain stage IV, which correlates with OH stage D, is not found in the North Amethyst area. Pyrites appear in both Bulldog Mountain stage V and North Amethyst stage 3.

Correlations between mineralogical stages of the North Amethyst vein system and mineralized segments of the southern part of the Amethyst vein are difficult because paragenetic relations for some of those areas are less well known. Two main stages of mineralization along the southern Amethyst vein system have been defined by Robinson (1981), and he has correlated the two assemblages with the A and B stages defined by Bethke and Rye (1979) for the OH vein. Minerals of southern Amethyst vein stage 2 correlate mineralogically with stage 1 of the North Amethyst vein (table 9). Galenas from the southern Amethyst stage-2 assemblage have lead-isotopic values that overlap the field defined by lead-isotopic compositions of galenas from the North Amethyst stage 1 (Foley, 1990).

Disseminated ore present in the hanging wall of the Amethyst fault at the southern end of the mineralized
structure, in the vicinity of an on-strike projection of the OH vein, has been examined by Guidice (1981). He has described two main hypogene stages and a supergene stage for disseminated minerals in volcanic rock located between the two veins. The first hypogene stage of mineralization may correlate with a combination of stages 1 and 2 of the North Amethyst vein and, as described by Plumlee (1989), with a combination of stages A through C of the OH vein and stages I through III of the Bulldog Mountain vein. This lumping of stages clearly reflects a lack of clear-cut paragenetic relations in the disseminated ore. The second hypogene stage for the disseminated ore correlates well with stage 3 of the North Amethyst vein.

The detailed paragenetic studies (table 9) demonstrate that base-metal minerals of the North Amethyst stage 1, OH and P vein stage B, and Bulldog Mountain stage II constitute the most widely distributed and persistent mineralogical stage, occurring in some form in all veins. Later stages of mineralization are more local and were interrupted by periods of dissolution. The North Amethyst vein retains the most complete record of mineralogical stages deposited anywhere in the district (table 9). In some of the other veins, only certain mineralogical stages are well developed, and many stages are absent or unrecognized. Thus, the North Amethyst vein probably has the most complete record of the changing chemical composition of the hydrothermal fluids.

**CONCLUSIONS**

1. The paragenesis of ore minerals of the North Amethyst vein system can be divided into two multistage associations separated by periods of intense brecciation. The earlier association has two stages, the first of which is characterized by abundant manganese minerals and the second by high gold content. The later association has three stages that consist primarily of base-metal sulfides, chlorite, quartz, carbonate minerals, and fluorite, in varying proportions.

2. Complete assemblages of the earlier Mn-Au stages of the North Amethyst system are not found elsewhere in the central San Juan region. In contrast, mineralogical stages and breccias identified in the base-metal-silica association can be correlated with paragenetic stages defined for veins of the central and southern Creede mining district.

3. The local and early occurrence of the Mn and Au assemblages may indicate that they either formed in a small hydrothermal cell that predated the extensive hydrothermal system from which ores of the Creede district are proposed to have been deposited (Bethke, 1988) or are remnants of a larger early hydrothermal system. If other early hydrothermal cells with similar characteristics were present in structures of the central and southern parts of the district, they may have been replaced or overprinted by later assemblages, or they remain to be discovered. Early-stage paragenetic mineral assemblages may hold the greatest potential for vein-related gold mineralization in Structures of the central caldera cluster of the San Juan Mountains.

4. The correlations of mineral assemblages described above have important implications for hydrologic modeling of the northern, central, and southern parts of the Creede district. The extensive correlation of base-metal-, silver-, and silica-rich assemblages suggests a similar source of fluids and metals for the ore mineralization and implies that the minerals were deposited in the veins by similar mechanisms and under similar conditions of ore formation.

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Coal Investigations Maps are geologic maps on topographic or planimetric bases at various scales showing bedrock or surficial geology, stratigraphy, and structural relations in certain coal-resource areas.

Oil and Gas Investigations Charts show stratigraphic information for certain oil and gas fields and other areas having petroleum potential.

Miscellaneous Field Studies Maps are multicolor or black-and-white maps on topographic or planimetric bases for quadrangle or irregular areas at various scales. Pre-1971 maps show bedrock geology in relation to specific mining or mineral-deposit problems; post-1971 maps are primarily black-and-white maps on various subjects such as environmental studies or wilderness mineral investigations.

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