

University of Nebraska - Lincoln

DigitalCommons@University of Nebraska - Lincoln

Geochemistry of Sulfate Minerals: A Tribute to
Robert O. Rye

US Geological Survey

2005

Alunite and the role of magmatic fluids in the Tambo high-sulfidation deposit, El Indio–Pascua belt, Chile

C.L. Deyell

University of British Columbia

R.O. Rye

U.S. Geological Survey

G.P. Landis

U.S. Geological Survey

T. Bissig

Queen's University, Kingston, Ontario, Canada

Follow this and additional works at: <https://digitalcommons.unl.edu/usgsrye>

 Part of the [Geochemistry Commons](#)

Deyell, C.L.; Rye, R.O.; Landis, G.P.; and Bissig, T., "Alunite and the role of magmatic fluids in the Tambo high-sulfidation deposit, El Indio–Pascua belt, Chile" (2005). *Geochemistry of Sulfate Minerals: A Tribute to Robert O. Rye*. 4.

<https://digitalcommons.unl.edu/usgsrye/4>

This Article is brought to you for free and open access by the US Geological Survey at DigitalCommons@University of Nebraska - Lincoln. It has been accepted for inclusion in Geochemistry of Sulfate Minerals: A Tribute to Robert O. Rye by an authorized administrator of DigitalCommons@University of Nebraska - Lincoln.

Alunite and the role of magmatic fluids in the Tambo high-sulfidation deposit, El Indio–Pascua belt, Chile

C.L. Deyell^{a,*}, R.O. Rye^b, G.P. Landis^b, T. Bissig^c

^a*Mineral Deposit Research Unit, Department of Earth and Ocean Sciences, University of British Columbia, 6339 Stores Rd., Vancouver, Canada V6T 1Z4*

^b*U.S. Geological Survey, MS-963 Denver Federal Center, Denver, CO, 80225, USA*

^c*Department of Geological Sciences and Geological Engineering, Queen's University, Kingston, Ontario, Canada K7L 3N6*

Accepted 1 June 2004

Abstract

The Tambo high-sulfidation deposit, located within the El Indio–Pascua belt in Chile, produced almost 25 t (0.8 M oz) of gold from altered Tertiary rhyodacitic volcanic rocks. Episodic magmatic-hydrothermal activity in the district occurred over at least 4 my and is characterized by several stages of acid-sulfate alteration, including magmatic-hydrothermal, magmatic steam, steam-heated, and apparent supergene assemblages. Two stages of Au±Ag mineralization are recognized and are hosted in barite and alunite within hydrothermal breccias and veins. Isotopic compositions of fluid in alunite show a dominant magmatic signature, with only a variable ¹⁸O-enriched meteoric water component throughout the entire hydrothermal process. Alunite ⁴⁰Ar/³⁹Ar ages constrain the timing of alteration and the duration of the hydrothermal system. Pre-ore alteration occurred at about 10 to 11 Ma and was contemporaneous with the volcanism of the Tambo Formation. Alunite from this stage of alteration occurs in the matrix of barren breccias and as fine intergrowths of alunite–quartz±clays that selectively replaced feldspars and pumice fragments. The textural relationships combined with stable-isotope systematics suggest a magmatic-hydrothermal origin for the alunite, with a local magmatic steam overprint. Early ore-stage alunite (8.7±0.2 Ma) occurs with barite and gold±walthierite within open spaces of the breccia matrix, and has δ³⁴S values (24–27‰) typical of magmatic-hydrothermal alunite, reflecting equilibrium between aqueous H₂S and SO₄²⁻. Fluid-inclusion ratios of H₂S/SO₂ (approximately 6) are consistent with ratios determined from stable-isotope data, and indicate reduced fluid conditions during ore deposition. Vapor-phase transport of Au, and deposition from condensed magmatic vapor rising from the brittle–ductile transition is inferred. Late gold coprecipitated with a third stage of alunite (8.2±0.2 Ma) that is characterized by nearly uniform chemical compositions and δ³⁴S values (1‰) similar to those for associated enargite, reflecting disequilibrium between H₂S and SO₄²⁻. This third-stage of alunite is isotopically and chemically similar to that of post-ore, coarse, banded alunite±hematite-quartz veins that crosscut the breccias in the Tambo area. Analyses of fluid-inclusion gas from alunite in these veins indicate high SO₂ and disequilibrium CO₂–CO–CH₄–H₂ species, consistent with a magmatic-steam origin. The gases are also depleted in He, and the late gold-bearing alunite was probably precipitated from rapidly ascending SO₂-rich vapors that were flashed from condensed magmatic

* Corresponding author. Present address: Centre for Ore Deposit Research, University of Tasmania, Private Bag 79, Hobart, Tasmania, 7001, Australia. Tel.: +61 3 6226 1735; fax: +61 3 6226 7662.

E-mail address: cdeyell@utas.edu.au (C.L. Deyell).

fluids located below the brittle–ductile transition. Alunite–huangite veins (8.6 ± 0.4 Ma) were formed intermediate to the two mineralizing events. Steam-heated alunite (8.9 ± 0.4 Ma), derived from the oxidation of H_2S within the vadose zone, is preserved at the highest elevations and overlaps with Stage 2 ore deposition. Post-mineral alunite \pm jarosite (7.3 ± 0.1 Ma) occurs as apparent supergene crosscutting veins and overgrowths. Stable-isotope data, however, indicate a strong magmatic component to both near-surface and late-stage alunite-forming fluids. This dominant magmatic fluid signature at high levels during the waning stages of hydrothermal activity is attributed to climatic and physiographic conditions specific to the El Indio–Pascua belt during the Middle to Late Miocene.

© 2004 Elsevier B.V. All rights reserved.

Keywords: Alunite; El Indio–Pascua belt; Tambo deposit; Acid-sulfate; Stable isotopes; Epithermal; Gold

1. Introduction

The Tambo high-sulfidation gold deposit is located in the El Indio–Pascua belt, within the Main Cordillera of Chile (Fig. 1). This belt hosts widespread zones of acid-sulfate hydrothermal alteration and several world-class epithermal deposits and prospects, including the El Indio mine (>15 t gold produced; Heather, personal communication, 2001), Pascua–Lama, with proven and probable resources of about 530 t (17 M oz) gold, and Veladero, with 317 t (11 M oz) gold (data from Barrick Gold Corporation website, 2003). The Tambo deposit is relatively small in comparison, and produced a total of just over 24.8 t (0.8 M oz) of gold from several tectonic-hydrothermal breccias and high-grade veins while it was in operation.

Previous studies of the El Indio district (Siddeley and Areneda, 1985; Bennet, 1995; Jannas, 1995; Jannas et al., 1999) have constrained the geological setting of the Tambo deposit. This region has many features in common with other high-sulfidation systems, although the deposit is unique in several aspects.

- (1) Limited erosion has allowed the preservation of many features of the hydrothermal system, including the near-paleosurface. As such, multiple stages of alunite [$KAl_3(SO_4)_2(OH)_6$] and acid-sulfate alteration can be recognized over several hundred meters of vertical extent. These stages include each of the four major types of acid-sulfate alteration described by Rye et al. (1992): magmatic-hydrothermal, magmatic-steam, steam-heated, and apparent supergene.
- (2) Mineralization is sulfide- and Cu-poor in comparison to that in most other deposits of this type (e.g., Cooke and Simmons, 2000;

Sillitoe, 1999). Ore is hosted primarily in barite and alunite \pm quartz of magmatic-hydrothermal origin. A second stage of Au mineralization is hosted in, and associated with, alunite of magmatic-steam origin (this study; Jannas et al., 1999).

In this paper, we examine shallow-level magmatic-hydrothermal processes within a well-constrained geological framework. The goal was to clarify the evolution of the Tambo deposit and, in particular, to determine the role of magmatic fluids throughout the life of the hydrothermal system, the mechanisms of ore-metal transport, and the nature of magmatic-steam processes. Emphasis is placed on the temporal and spatial relationship between alunite and Au mineralization because the isotopic, geochronological, and geochemical characteristics of alunite can be used to resolve the nature of source fluids (e.g., Bethke, 1984; Rye et al., 1992). Multiple stages of alunite have been recognized at the Tambo deposit. The relative timing of major alteration and mineralizing events was determined from the paragenetic sequence defined by Jannas (1995), with additional constraints from field relations and new $^{40}Ar/^{39}Ar$ ages for alunite.

2. Geological framework

2.1. Regional geology

The El Indio–Pascua belt is in the center of a presently amagmatic segment of the Andean Cordillera. The lack of recent volcanism is attributed to subhorizontal subduction of the Nazca plate beneath the South American continent between $27^\circ 30'$ and

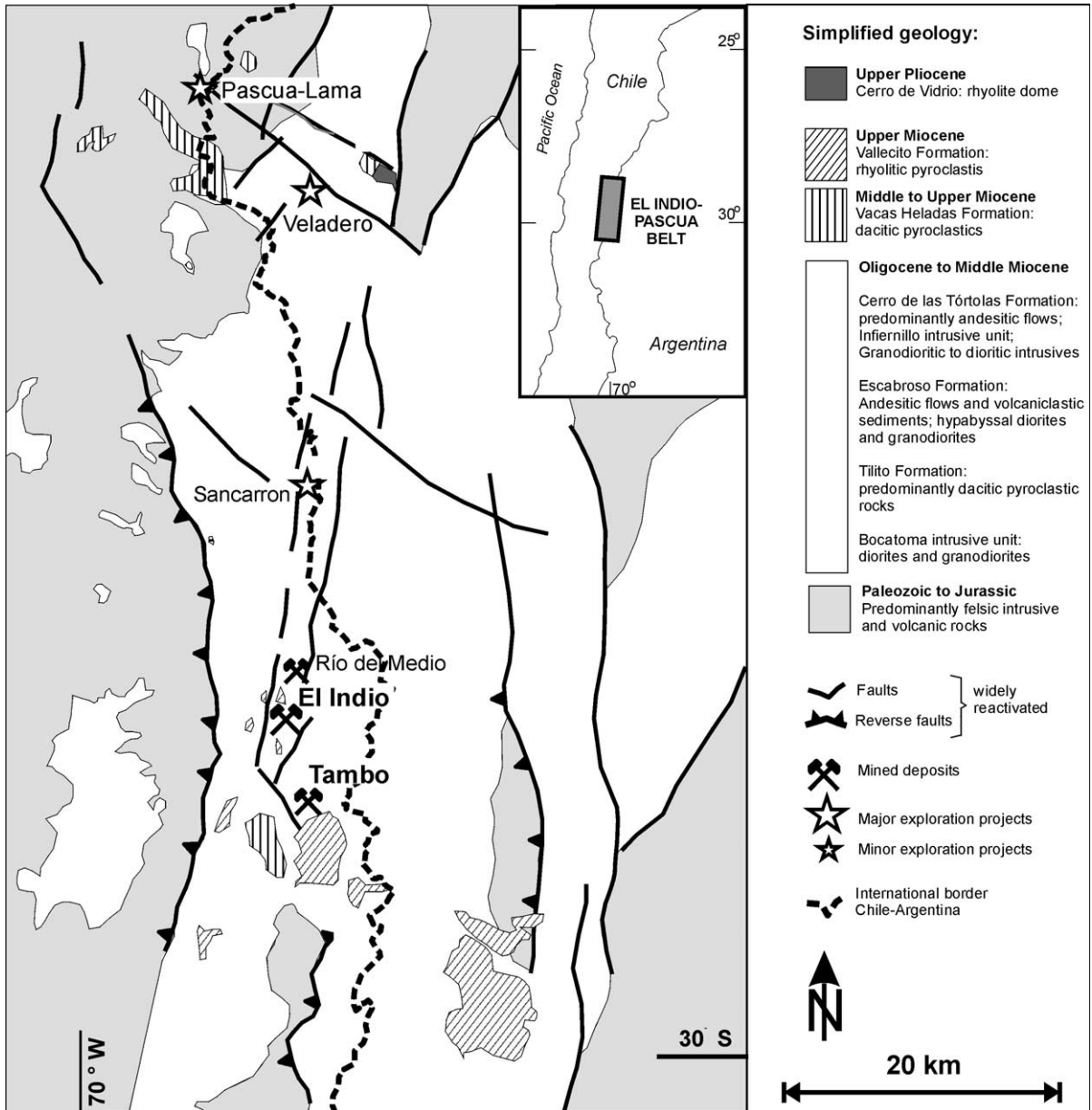


Fig. 1. Simplified geology of the El Indio–Pascua belt, showing the locations of the Tambo and El Indio deposits.

33°S (Barazangi and Isacks, 1976). The district is approximately 100 km long and 30–40 km wide, and straddles the Chile–Argentina border (Fig. 1). Hydrothermal systems formed within a NNE-striking tectonic depression bounded by the steeply west-dipping high-angle reverse Baños del Toro Fault to the west and opposing structures in the Valle del

Cura region of Argentina to the east. Within this block, an Upper Paleozoic to lower Jurassic basement consisting predominantly of calc-alkaline felsic intrusive suites and volcanic-sedimentary rocks (Martin et al., 1999) is overlain by up to 1500 m of Tertiary subaerial volcano-sedimentary strata. The latter are extensively preserved in the southern part

of the belt, where the Tambo and El Indio mines are located (Fig. 1), but are less extensive in the northern extremity of the district near Pascua–Lama and Veladero (Maksaev et al., 1984; Martin et al., 1995).

The Tertiary volcanic sequence is briefly summarized here (Martin et al., 1995; Bissig, 2001). Up to 1200 m of dacitic and rhyodacitic tuffs with volcanoclastic sediments and subordinate basaltic and andesitic flows comprise the 23–27-Ma Tilito Formation. It is separated from the overlying 17.5–21-Ma Escabroso Formation by an angular unconformity marked by a locally well-developed regolith (Heather, personal communication, 2001). The Escabroso Formation

consists predominantly of andesitic lavas, hypabyssal intrusive bodies, and volcanoclastic sedimentary rocks. After a brief deformation event, igneous activity resumed and is represented by the 14–17-Ma andesitic Cerro de las Tórtolas Formation and the contemporaneous Infiernillo intrusive unit. Dacitic pyroclastic rocks of the Tambo Formation (formerly the Vacas Heladas Formation; Martin et al., 1997) erupted from isolated centers between 11 and 12.7 Ma. A single 7.8-Ma dacite dike, the only igneous rock coeval with mineralization in the district, has been reported to occur at Pascua (Bissig, 2001). The 5.5–6.2-Ma Vallecito Formation rhyolitic tuffs are restricted to an ignimbrite

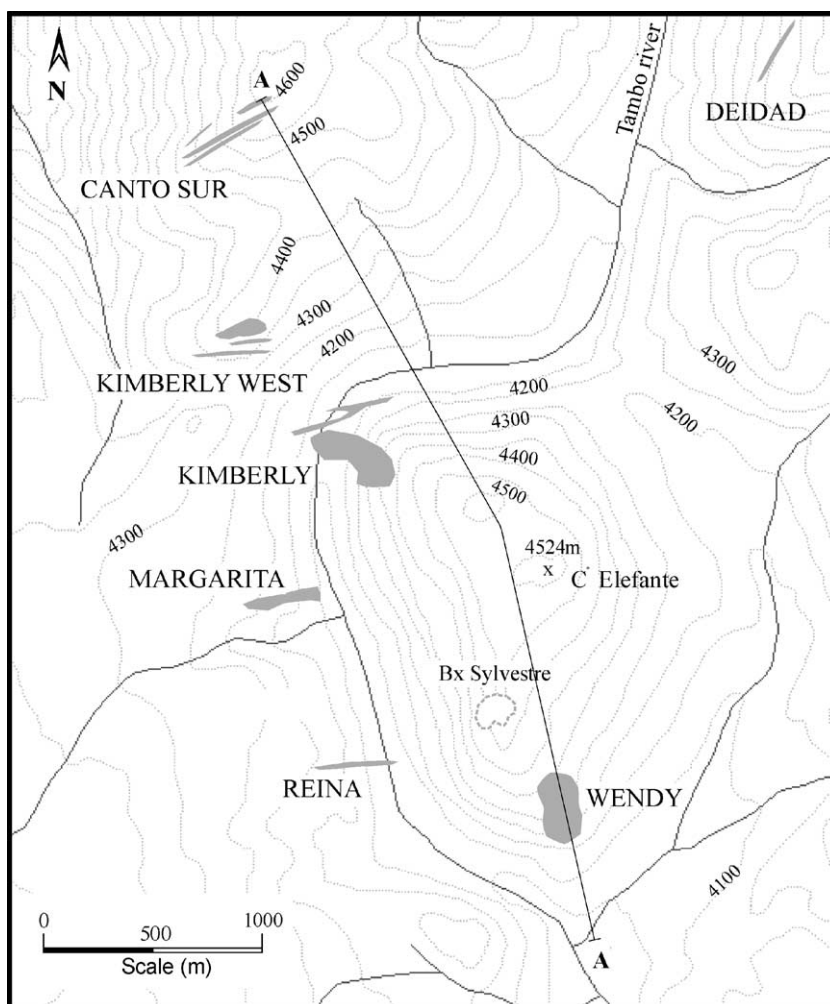


Fig. 2. Map of the Tambo property (modified from Jannas et al., 1999). Shaded areas indicate zones of mineralization.

sheet in the Valle del Cura (Ramos et al., 1989) and other, isolated occurrences in the El Indio–Tambo area. Volcanism ceased in the Upper Pliocene after eruption of the 2-Ma Cerro de Vidrio rhyolite dome in the northeastern corner of the belt (Bissig, 2001).

2.2. Local geology and mineralization

The geology of the El Indio–Tambo area is dominated by an upper Oligocene to Upper Miocene sequence of intensely fractured volcanic and volcanoclastic rocks. The epithermal mineralization is hosted by rhyolitic to dacitic, lithic ash-flow tuffs of the Tilito Formation (Heather and Diaz, 2000; Bissig, 2001) and, at El Indio, is overlain by andesitic lavas of the Escabroso Formation. Intrusive units and hydrothermal alteration of Escabroso and Infiernillo age are also present in the area, but $^{40}\text{Ar}/^{39}\text{Ar}$ ages for gold mineralization both at El Indio and Tambo are significantly younger than these intrusions (Bissig, 2001).

Gold (and Ag) mineralization at Tambo is hosted in three major hydrothermal breccias and several high-grade veins. The geology of this deposit is summarized after Siddeley and Areneda (1985) and Jannas et al. (1999). Two of the major breccia deposits, Kimberly and Wendy, are on the northwest and south flanks of Cerro Elefante, respectively (Figs. 2 and 3). These deposits have lateral and vertical dimensions of up to 300 and 400 m, respectively, and consist of matrix-supported, monolithic breccias that contain

altered fragments of volcanic tuffs (Jannas et al., 1999). The main ore zone occurs between 4200 and 4000 m a.s.l. (all elevations are reported as elevation above sea level) with grades up to $4\text{--}10\text{ g t}^{-1}$ Au and 10 g t^{-1} Ag (Jannas et al., 1999). The Canto Sur deposit is at a higher elevation on Cerro Canto, which separates the El Indio and Tambo mines. All three breccias are located at, or close to, the intersection of north–northwest and east–northeast fault systems in areas of intense shears and extensional joints (Jannas et al., 1999). Vein mineralization at Tambo is primarily hosted by the Reina, Deidad, and Indigena structures (Fig. 2). These veins are typically narrow (1–4 m wide), of limited vertical extent (<200 m), and consist primarily of barite and alunite \pm quartz. Ore grades averaged 15 g t^{-1} Au and 30 g t^{-1} Ag, but locally exceeded 1000 g t^{-1} Au (Jannas et al., 1999).

Two stages of mineralization were recognized by Jannas et al. (1999). Early Au mineralization (Table 1) is closely associated with barite in both the veins and the breccia deposits. Gold and argentian gold, minor sulfides (including enargite, galena, and pyrite), and native tellurium and other Te minerals are concentrated along growth planes in the barite. Trace amounts of sulfides and gold occur interstitial to alunite overgrowths on barite grains. Gold also occurs locally on the rims of the breccia fragments. The second stage of mineralization is characterized by native gold that occurs in fractures and vugs, and interstitial to grains of alunite, hematite, jarosite, and barite. Inclusions of

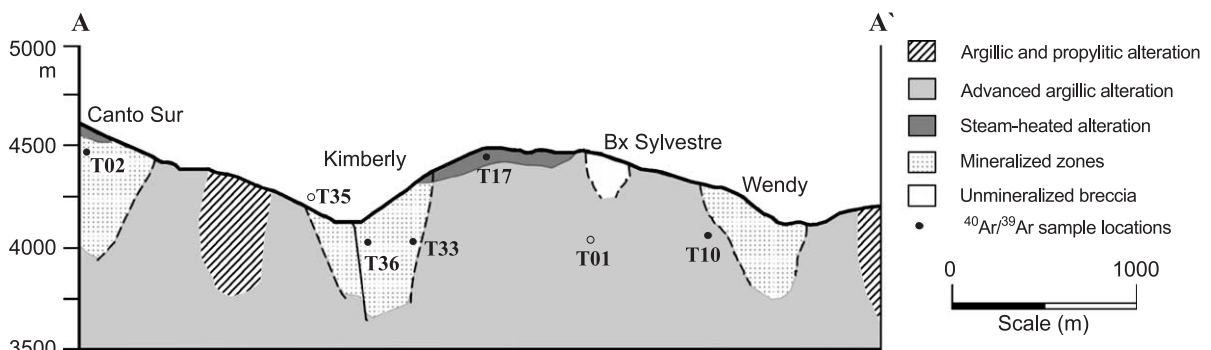


Fig. 3. Schematic section showing a projection of the three major mineralized breccias (Canto Sur, Kimberly, and Wendy) and zones of alteration at Tambo. Line of section is marked in Fig. 2. Also shown are locations of $^{40}\text{Ar}/^{39}\text{Ar}$ samples; open symbols indicate samples projected onto the section.

Table 1
Paragenetic sequence and characteristics of the eight different stages of alunite formation at Tambo^a

	Brecha Sylvestre	Stage 1	Stage 2	Stage 3	Banded veins	Steam-heated	Huangite-bearing veins	Late-stage oxidation
⁴⁰ Ar/ ³⁹ Ar age	10.42±0.28	11.00±0.27	8.71±0.21	8.24±0.15	–	8.86±0.38	8.63±0.41	7.25±0.14
Alunite occurrence	Matrix fill; with qtz and late jar	Qtz–alun±kao, dick, pyl alteration of Amiga tuff	Vein and bx matrix alun associated with bar, wal, trace qtz, and late jar	Fine-grained to coarse, breccia and vein filling alun	Coarse, banded alun veins with hem	Surficial qtz–kao– sulfur±alun alteration	Fine-grained, thin pink alun–huag veinlets with trace kao	Fine-grained alun with minor qtz; late jar
Mineralization	–	–	Gold, minor sulfides (en, gn, py), native Te, Te–Au minerals, native gold	Native gold, trace en	–	–	–	–

Abbreviated mineral names; alunite (alun), quartz (qtz), kaolinite (kao), dickite (dick), pyrophyllite (pyl), walthierite (wal), jarosite (jar), enargite (en), galena (gn), pyrite (py), hematite (hem), huangite (huag). Reported ⁴⁰Ar/³⁹Ar ages in bold are from this study; other ages are reported in Bissig (2001).

^a Based on the nomenclature of Jannas et al. (1999) with additions from this study.

native gold also occur in alunite. Sulfides are rare, although minor enargite occurs at depth in the Wendy breccia, intergrown with coarse grains of alunite. Early tellurides typically are replaced by minerals such as poughite, rodalquilarite, and tellurite (Jannas et al., 1999).

3. Methods of analysis

Alteration assemblages were identified by field observations and detailed petrography. At the time of this study, access was limited to surficial outcrops, open pits (Kimberly, Wendy, and Canto Sur), and drill core. The mineralogy and character of each paragenetic stage were determined by examination of thin and polished sections using standard optical methods and X-ray diffraction (XRD). In addition, short-wave infrared (SWIR) analyses were made in situ using a Portable Infrared Mineral Analyzer (PIMA), which is a field-portable spectrometer. Spectra were analyzed using PIMAVIEW 3.0 software, and mineral identification was made by comparison to standards in the SPECMIN database (Thompson et al., 1999). Repre-

sentative samples from each paragenetic stage were selected for isotopic and geochemical analyses.

Six alunite samples were dated by ⁴⁰Ar/³⁹Ar laser step-heating methods. All analyses were done with a Mass Analyzer Products MAP 216 mass spectrometer at the Queen's University ⁴⁰Ar–³⁹Ar laboratory. About 10 mg of each sample was irradiated for 7.5 h at the McMaster University nuclear reactor in Hamilton, Canada, using biotite standard Mac-83 as a radiation-flux monitor (24.36±0.17 Ma; Sandeman et al., 1999). *J*-values for individual samples were determined by second-order polynomial interpolation and, for the samples discussed in this article, were about 2.4×10^{−3} and varied by less than 0.02×10^{−3}. Samples were step-heated using a defocused LEXEL 3500 Ar laser-beam. Ages were calculated using the decay constants suggested by Steiger and Jäger (1977), and all errors are given at 2σ.

Geochemical characteristics of alunite were determined by analysis of individual grains and sample composites. Grain mounts and polished sections were analyzed by scanning electron microscopy with energy-dispersion spectroscopy (SEM-EDS) and electron-probe microanalysis (EPMA). EPMA data

were collected on a fully automated CAMECA SX-50 microprobe operated in the wavelength-dispersion mode and at 15 kV excitation voltage, 10 nA beam current, 15 μm beam diameter, and total count time <65 s. Analytical parameters were chosen to minimize beam damage caused by the volatilization of alkali elements (Petersen and Thompson, 1992). Data reduction was done with the PAP method (Pouchou and Pichoir, 1985). Alunite separates (~1 g samples) were carefully hand picked and treated using a 1:1 HF–H₂O solution to remove silicate contaminants (Wasserman et al., 1992). The purity of separates was checked by XRD, and trace to minor amounts (i.e., ≤ 10 vol.%) of quartz, clays, or other impurities may have been present. Samples were analyzed by inductively coupled plasma mass spectrometry (ICP-MS) for trace-metal and rare-earth-element (REE) contents by ALS Chemex Laboratories, Canada. For trace-metal analysis, samples were dissolved in a mixture of HClO₄, HNO₃, and HF acids to ensure total dissolution. For REE determination, samples were fused with lithium metaborate prior to analysis. Data for REE in alunite are normalized to the average REE content of volcanic tuffs from the Tilito Formation (Bissig, unpublished data; average values given in Table 2) that host ore in the El Indio district.

Complete stable isotopic analyses ($\delta^{34}\text{S}$, δD , $\delta^{18}\text{O}_{\text{SO}_4}$, and $\delta^{18}\text{O}_{\text{OH}}$) were obtained for alunite from each paragenetic stage. Sulfur-isotope data

were also obtained for selected samples of barite and enargite. All data were collected at the U.S. Geological Survey Isotope Laboratory in Denver, USA. Samples were separated and prepared according to the procedure of Wasserman et al. (1992). Sulfur isotopic ratios were determined by an on-line method using an elemental analyzer coupled to a Micromass Optima mass spectrometer (Gieseman et al., 1994). Analytical precision is better than $\pm 0.2\%$. D/H analyses were performed by a step-heated technique modified from Godfrey (1962), with analytical precision better than $\pm 3\%$ (Wasserman et al., 1992). Oxygen-isotope data for alunite were collected for both sulfate and hydroxyl oxygen. For analysis of $\delta^{18}\text{O}_{\text{SO}_4}$, alunite was dissolved in a hot NaOH solution, and sulfate was precipitated as BaSO₄ (Wasserman et al., 1992). The BaSO₄ precipitate was then analyzed by fluorination with BrF₅ at 580 °C in accordance with the standard analytical procedure of Clayton and Mayeda (1963), and as used by Pickthorn and O’Neil (1985). Analytical precision is estimated at $\pm 0.15\%$. Values of $\delta^{18}\text{O}_{\text{OH}}$ were determined by material balance using the total-O isotopic composition of alunite (measured by fluorination with BrF₅, described above) and the $\delta^{18}\text{O}_{\text{SO}_4}$ results. Several oxygen-isotope analyses were also determined by pyrolysis with a Finnigan TC/EA coupled to a Finnigan Delta Plus XL mass spectrometer using continuous-flow methods modified from Kornexl et al. (1999). Analytical precision is better than $\pm 0.3\%$.

Analyses of fluid-inclusion gas were done at the U.S. Geological Survey fluid-inclusion and noble-gas laboratories in Denver, USA. All samples were initially crushed in a sealed stainless steel tube to release gas from the inclusions. After analysis of this gas, the crushed alunite was heated at 200 °C for 1 h in the same tube to generate thermally released gas. The temperature chosen was sufficient to decrepitate fluid inclusions (especially vapor-rich inclusions) that still remained after crushing, yet avoided thermal decomposition of the alunite. The released gases from both extractions were split for gas-composition analysis on a calibrated quadrupole mass spectrometer (Pfeiffer Vacuum Prisma), and the remaining gas was analyzed for He and Ne isotopes on a high-resolution MAP 215-50 noble-gas instrument. Gas species were analyzed at approximately 10^{-8} Torr, with each

Table 2
Average major and REE geochemical data for unaltered rocks of the Tilito Fm ($n=3$)

	wt.%		ppm
Na ₂ O	2.89	La	31.6
K ₂ O	4.24	Ce	62.6
CaO	4.35	Pr	7.2
MgO	1.59	Nd	25.2
MnO	0.09	Sm	4.8
Fe ₂ O ₃	5.05	Eu	1.0
Al ₂ O ₃	16.39	Gd	3.9
P ₂ O ₅	0.16	Tb	0.6
SiO ₂	62.12	Dy	3.5
TiO ₂	0.66	Ho	0.7
Total	97.54	Er	2.1
		Tm	0.3
		Yb	2.1
		Lu	0.3

Data from Bissig (2001); Fe₂O₃ from total Fe. REE by ICP-MS.

analysis consisting of an average of 40 to 60 matrix calculations that correct for ion sensitivity and gas ionization fragmentation, and which calibrate the intensity response to standards. Results were determined as mol% summed to 100% when normalized to nitrogen. The remaining gas was then processed to extract He and Ne by cryogenically removing other gases on charcoal, gettering the gas (Saes ST707), and concentrating the purified He and Ne in a small-volume charcoal trap. Helium, and then Ne, were separately analyzed by selective thermal desorption from the charcoal and expansion into the mass spectrometer. Helium and Ne calibration was achieved by processing laboratory air corrected for barometric pressure, temperature, and relative humidity using identical procedures ($^4\text{He}=6.414\text{e}^{-12}$ mol $\text{V}^{-1}\pm 0.016\%$, $^3\text{He}=9.951\text{e}^{-20}$ mol $\text{cps}^{-1}\pm 0.144\%$, $^3\text{He}/^4\text{He}$ detector discrimination= $1.03074\pm 1.68\%$; Ne isotopes are treated as normalized percentages and ratios only, but are corrected for $^{40}\text{Ar}^+$ and CO_2^+ isobaric interferences to ^{20}Ne and ^{22}Ne).

4. Alunite paragenesis in the Tambo system

Multiple stages of alunite deposition are recognized in the Tambo district. The paragenetic sequence (Table 1) is adapted from Jannas et al. (1999) based on new field and petrographic observations, and reliable alunite $^{40}\text{Ar}/^{39}\text{Ar}$ ages from this study (see Section 5) and Bissig (2001). The different occurrences of alunite are described below.

4.1. Brecha Sylvestre

Brecha Sylvestre is a relatively small, barren breccia body at the southwestern end of Cerro Elefante (Fig. 2). The breccia occurs between 4430

and 4480 m, approximately 200 m above the zones of highest gold production in the Wendy and Kimberly orebodies. The breccia is mainly clast-supported, locally silicified, and shows evidence of intense acid-leaching that removed most of the matrix material (Bennet, 1995). The matrix consists of alunite and minor quartz that line vugs (Fig. 4a). Alunite typically occurs as pale white to yellow, bladed to acicular crystals that are 20 to 200 μm in length. Late jarosite is also common and overgrows silicified breccia fragments and matrix crystals.

4.2. Stage 1—pre-ore acid-sulfate alteration

Widespread acid-sulfate alteration of the Tilito Formation wallrock is recognized throughout the Tambo area (Stage 1). This assemblage is surrounded by argillic- and propylitic-altered rocks away from the main orebodies (this study; Jannas, 1995). The occurrence of altered clasts with silicified rims in the hydrothermal breccias indicates that alteration occurred prior to brecciation and subsequent mineralization (Jannas et al., 1999). Alteration occurs as fine intergrowths of alunite-quartz that selectively replaced feldspar crystals and locally formed massive clusters in the matrix of the wallrock (Fig. 4b). Pumice clasts within fragments of dacite tuff in the breccia pipes are also replaced (Jannas, 1995). Original rock textures are generally preserved. Associated minerals include clays (kaolinite–dickite), pyrophyllite, rare pyrite, and diaspore. Alunite commonly occurs as clusters of tabular to bladed crystals between 50 μm and 2 mm in length. Crystals locally have cores of aluminum-phospho-sulfate (APS) minerals, including woodhouseite, svanbergite, and florencite (Table 3). Alunite grains are also locally pitted, have corroded margins, and are rarely overprinted by fine-grained (10–20 μm) alunite \pm quartz.

Fig. 4. (a) Medium-grained, tabular alunite crystals in the matrix of Brecha Sylvestre (crossed polars). Scale bar=50 μm . (b) SEM backscattered-electron (BSE) image of Stage 1 quartz (qtz)–alunite (aln) alteration. Bright zones within alunite grains are inclusions of woodhouseite–svanbergite. Scale bar=25 μm . (c) BSE image of walthierite (light gray)–alunite (dark gray) zoning in Stage 2 alunite from the Canto Sur deposit. Alunite is hosted in barite (white). Scale bar=50 μm . (d) BSE image of native gold (white) associated with Stage 2 alunite (gray) from Wendy (DDH-92a, 206.7 m). Circle in centre of image is from electron-beam damage. Scale bar=25 μm . (e) Fine-grained (white) Stage 3 alunite (aln) overgrown on coarser-grained Stage 3 alunite (gray) in the matrix of the Wendy breccia. Minor enargite (en) is associated with the coarse-grained alunite. Scale bar=1 cm. (f) Banded alunite–hematite vein from the Kimberly deposit. Scale bar=1 cm. (g) BSE image of zoned alunite (light gray)–huangite (darker gray)–woodhouseite–svanbergite (white) grains from the Kimberly pit. Scale bar=10 μm . (h) BSE image of fine-grained, fspseudocubic grains of alunite (gray) with minor jarosite (white). Scale bar=10 μm .

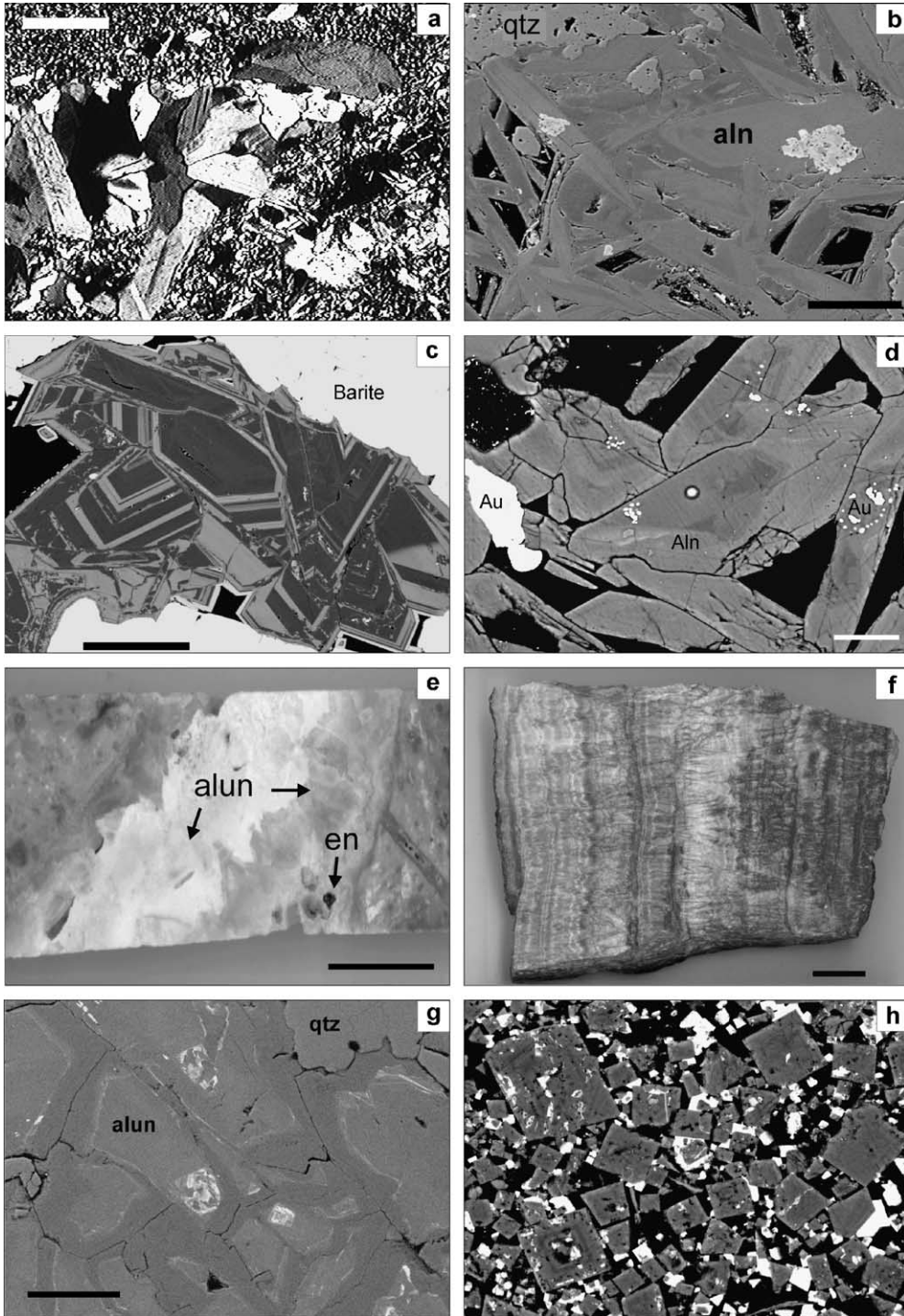


Table 3
Minerals of the alunite–jarosite family identified in the Tambo area

Alunite–jarosite group	alunite	$\text{KAl}_3(\text{SO}_4)_2(\text{OH})_6$
	natroalunite	$\text{NaAl}_3(\text{SO}_4)_2(\text{OH})_6$
	minamiite	$(\text{Na},\text{K},\text{Ca})_2\text{Al}_6(\text{SO}_4)_4(\text{OH},\text{H}_2\text{O})_{12}$
	huangite	$\text{CaAl}_6(\text{SO}_4)_4(\text{OH})_{12}$
	walthierite	$\text{BaAl}_6(\text{SO}_4)_4(\text{OH})_{12}$
jarosite	$\text{KFe}_3(\text{SO}_4)_2(\text{OH})_6$	
Beudantite group	svanbergite	$\text{SrAl}_3[(\text{P},\text{S})\text{O}_4]_2(\text{OH},\text{H}_2\text{O})_6$
	woodhouseite	$\text{CaAl}_3[(\text{P},\text{S})\text{O}_4]_2(\text{OH},\text{H}_2\text{O})_6$
Crandallite group	crandallite	$\text{CaAl}_3(\text{PO}_4)_2(\text{OH},\text{H}_2\text{O})_6$
	florencite-(Ce)	$\text{CeAl}_3(\text{PO}_4)_2(\text{OH},\text{H}_2\text{O})_6$
	florencite-(La)	$\text{LaAl}_3(\text{PO}_4)_2(\text{OH},\text{H}_2\text{O})_6$

4.3. Stage 2—early Au mineralization

Stage 2 alunite post-dates the emplacement of the three major breccias and is closely associated with early deposition of Au (Jannas et al., 1999). Alunite typically occurs as euhedral, golden yellow, bladed crystals up to 1 cm long that have overgrown barite and breccia fragments. Finer-grained tabular crystals, <50 μm in length, also occur. Alunite of this stage is recognized by complex compositional zoning and the presence of walthierite (the Ba analogue of alunite; Table 3). The type of locality for walthierite is the Reina vein (Beane, 1991; Li et al., 1992), where the mineral forms irregular to oscillatory compositional zones (Fig. 4c) and aggregates in coarse-grained alunite crystals. Stage 2 alunite also occurs in the Kimberly and Canto Sur breccias in close association with barite and traces of quartz.

4.4. Stage 3—late Au mineralization

Deposition of Stage 3 alunite overlapped with the later stage of Au deposition and continued after mineralization ceased. Alunite is characterized by its nearly stoichiometric composition and variable crystal size and shape. Fine-grained alunite occurs as lath-like to thin bladed crystals, rarely >30 μm long, which form overgrowths on breccia fragments and Stage 2 barite crystals in the Kimberly and Canto Sur breccia pipes. Alunite commonly exhibits the characteristic ‘earthy’ texture described by Jannas et al. (1999) and locally contains inclusions of native gold (Fig. 4d). Coarser-grained varieties (up to 2 cm) of white to pinkish brown, bladed alunite occur in veins, as cement to breccia fragments, as overgrowths on Stage 2 alunite crystals, and as local intergrowths with

enargite (Fig. 4e). Fine-grained alunite crystals also post-date the coarser bladed veins and breccia matrix. Minor amounts of quartz and kaolinite are associated with this stage. Late jarosite and scorodite [$\text{FeAsO}_4 \cdot 2\text{H}_2\text{O}$] that overprint Stage 3 alunite are common.

4.5. Veins of coarse banded alunite (\pm hematite)

Veins, up to 20 cm wide, of coarsely crystalline alunite \pm hematite (Fig. 4f) are common in the upper and marginal parts of the Kimberly open pit, and occur locally in the Wendy breccia. The veins crosscut both Stage 1 alteration and hydrothermal breccias, but the timing of vein formation relative to Stage 3 alunite is unknown. The veins are characterized by coarse alunite crystals, 1 to 5 mm long, that grew in successive bands outward from the wallrock. Minor hematite and rare quartz \pm jarosite occur interstitial to the alunite crystals. The presence of hematite gives the veins a distinct reddish color that allows easy distinction from the surrounding wallrock. Where hematite is absent, the veins have a cream-white color.

4.6. Huangite-bearing veins

Another variety of late veins occurs in the Kimberly and Kimberly West open pits. These veinlets are thin, pink to white, cryptocrystalline, and are composed of alunite, huangite (the Ca analogue of alunite; Table 3) \pm kaolinite, quartz, and trace pyrite (Fig. 4g). Alunite occurs as clusters of tabular to anhedral grains averaging 50 to 100 μm in length, locally overprinted by late jarosite and scorodite. The veinlets cut Stage 1 alteration, but timing with respect to Stage 2 or later events is unknown. Huangite had not been recognized previously in the Tambo area, but the type locality is in the El Indio district, where the mineral occurs in association with kaolinite and pyrite (\pm alunite, pyrophyllite, sericite, minamiite, and woodhouseite) adjacent to the El Indio Campana B vein (Li et al., 1992). Alunite in this vein has been dated at 6.21 ± 0.26 Ma and represents the youngest alteration event in the district (Bissig, 2001).

4.7. Near-surface alteration

A powdery, friable assemblage of silica–kaolinite \pm alunite and native sulfur is present at higher elevations

(4300–4500 m) in the Tambo district and occurs as alteration blankets on top of Cerro Elefante, Cerro Canto, and the adjacent Azufreras and Sol Poniente areas. Alteration is commonly pervasive, with the matrix of the host rocks and feldspars in tuff fragments completely replaced. Alteration also extends downward along fractures, particularly beneath the surficial alteration zone on Cerro Elefante, and locally overprints older acid-sulfate assemblages. Alunite in the near-surface zone is restricted to irregular pods and lenses and occurs as fine-grained (<15–20 μm), tabular, and less commonly pseudocubic crystals intergrown with fine-grained silica. The mineralogy and distribution of this assemblage is typical of steam-heated alteration observed in geothermal fields (e.g., Schoen et al., 1974; Rye et al., 1992).

4.8. Late alunite veins \pm jarosite

Late, fine-grained to cryptocrystalline veinlets of alunite \pm jarosite are common in the Tambo area. These veinlets occur at all depths investigated by drilling (down to 3800 m elevation) and crosscut all other alteration types. The alunite is typically fine- to medium-grained (10–50 μm), tabular to bladed in habit, and is locally intermixed with quartz. Alunite is commonly overgrown and locally crosscut by fine-grained jarosite. In rare occurrences, extremely fine-grained (2–10 μm), pseudocubic alunite is intergrown with kaolinite \pm jarosite (Fig. 4h) in veinlets that crosscut wallrock altered by illite–smectite \pm chlorite. These late-stage veins were previously interpreted to be supergene in origin (Guerra, personal communication) on the basis of their distribution and field characteristics.

5. Geochronology

Six alunite samples were dated by the $^{40}\text{Ar}/^{39}\text{Ar}$ laser step-heating method to supplement existing age data for the Tambo area (Jannas et al., 1999; Bissig, 2001) and to confirm the relative timing of the alteration and mineralizing events. Results (Table 4) indicate that Stage 1 wallrock alteration (11.0 ± 0.3 Ma) occurred 2 to 3 my prior to either Stage 2 or Stage 3 ore deposition. Early alteration is contemporaneous with Tambo Formation volcanism, and extensive hydrothermal alteration of this age can be traced throughout the El Indio–Pascua belt (Bissig, 2001). A sample of coarse, vug-fill alunite from Brecha Sylvestre, dated at 10.4 ± 0.3 Ma (Bissig, 2001), is also pre-ore and may overlap with Stage 1 alteration.

The timing of Stage 2 mineralization is constrained by $^{40}\text{Ar}/^{39}\text{Ar}$ data (8.7 ± 0.2 Ma) for a sample of coarse-grained alunite from the Reina vein that contains inclusions of gold. Alunite in this sample is also intergrown with barite that contains precious metals. Stage 3 mineralization is slightly younger according to data for medium-grained alunite (8.2 ± 0.2 Ma) sampled from the matrix of the Kimberly breccia.

The near-surface, blanket style of alteration (8.9 ± 0.4 Ma) was sampled at Cerro Elefante, above the Kimberly deposit. Alteration overlaps in time with Stage 2 and may represent the surficial (steam-heated?) expression of early Au mineralization.

The timing of huangite-bearing veins (8.6 ± 0.4 Ma) relative to mineralization is uncertain. The relatively large error for this analysis is attributed to the small grain size of alunite and a smaller quantity of Ar gas released during heating. These veins may overlap with either Stage 2 or Stage 3, or may be

Table 4
 $^{40}\text{Ar}/^{39}\text{Ar}$ results for six stages of Tambo alunite

Stage	Sample	Lab no.	Integrated date (Ma $\pm 2\sigma$)	Plateau age (Ma $\pm 2\sigma$)	% ^{39}Ar in the plateau	Correlation date (Ma $\pm 2\sigma$)	% ^{39}Ar in the correlation	MSWD	Initial $^{40}\text{Ar}/^{39}\text{Ar}$ ($\pm 2\sigma$)
1	T10a	TB-106	10.87 (0.34)	11.00 (0.27)	94.4	11.10 (1.46)	100.0	0.16	291.17 (92.11)
2	T01	TB-110	8.71 (0.21)	–	–	8.72 (0.63)	100.0	0.43	290.79 (166.77)
Ca-vein	T35	TB-103	8.61 (0.45)	8.63 (0.41)	87.2	6.97 (6.28)	100.0	3.70	526.85 (1419.61)
SH	T17	TB-101	8.68 (0.59)	8.86 (0.38)	91.7	8.97 (3.45)	100.0	0.02	293.77 (183.44)
LV	T02a	TB-104	7.18 (0.15)	7.25 (0.14)	88.4	7.27 (0.13)	100.0	0.52	226.54 (247.77)
BV	T33	TB-111	7.38 (2.88)	7.70 (2.80)	90.3	8.15 (12.78)	100.0	0.04	293.76 (672.32)

Ca-vein=huangite-bearing vein, SH=steam-heated, LV=late vein, BV=banded vein. Calculated from data given in Deyell (2001).

intermediate between the two. Crosscutting relations could not be determined in the field. It is clear, however, that alunite in these veins is unrelated to the much younger huangite-bearing Campana B vein at El Indio (6.2 ± 0.3 Ma; Bissig, 2001).

Late-stage alunite (\pm jarosite) was sampled from thin veinlets that crosscut Stage 1 alteration in the Canto Sur deposit. The plateau age for this sample (7.3 ± 0.1 Ma) is approximately 1 my younger than the main mineralizing events in the Kimberly and Wendy areas. It should be noted, however, that Bissig (2001) reported an age of 7.1 ± 0.2 Ma for alunite thought to be related to Au deposition in the Canto Sur breccia. The paragenesis of this sample is unclear. The style and mineralogy of both the alteration and the ore assemblages at Canto Sur are similar to those in other parts of the Tambo deposit. These features suggest a common origin. If mineralization in this area is younger, however, then late-stage alteration is coeval with, and not younger than, Canto Sur ore-stage alunite.

Repeated attempts to obtain accurate age data for banded-vein alunite were unsuccessful because the samples failed to release sufficient Ar. The reported plateau age of 7.7 ± 2.8 Ma does not help constrain the timing of the banded veins relative to other stages of Tambo alteration and gold deposition.

6. Stable-isotope study

Stable-isotope analyses of alunite were determined for each paragenetic stage. Alunite contains four stable-isotope sites and $\delta^{34}\text{S}$, δD , $\delta^{18}\text{O}_{\text{SO}_4}$, and $\delta^{18}\text{O}_{\text{OH}}$ data can be used to characterize the fluid source and environment of deposition for each stage of alteration (Rye et al., 1992). Brief sample descriptions and isotope results are presented in Table 5. The data show considerable variation in $\delta^{34}\text{S}$ of alunite, with values ranging from 0‰ to 27‰ (Fig. 5). Limited data for Stage 3 enargite range from about -2% to -4% , consistent with values reported in Jannas et al. (1999).

Estimation of $\delta^{34}\text{S}$ for bulk S in the magma (1–3‰; Fig. 5) is based on data for magmatic-steam alunite (cf. Rye et al., 1992) and agrees with estimates presented by Jannas et al. (1999). The δD of ambient meteoric waters is estimated at $-100 \pm 10\%$ (Taylor, personal communication). However, isotope data for

jarosite of supergene origin from the Pascua deposit (Deyell, 2001), located to the north of El Indio–Tambo, suggest that this value has been slightly lower (about -125%). Variations of this magnitude are expected during the 3–4 my duration of the Tambo hydrothermal system, and a range of meteoric δD values is shown on all isotope plots (Figs. 6–8) accompanying the following discussions.

Stable-isotope systematics for each paragenetic stage are presented below. For ease of data interpretation and discussion, some samples are grouped with other, related stages. In each section, $\delta^{34}\text{S}$ data are discussed first, followed by $\delta^{18}\text{O}$ and δD relations. The $\delta\text{D}_{\text{H}_2\text{O}}$ and $\delta^{18}\text{O}_{\text{H}_2\text{O}}$ values of fluids in equilibrium with each alunite sample are calculated from the fractionation equations of Stoffregen et al. (1994) over a range of temperatures. These temperatures are estimated from textural data, $\Delta^{18}\text{O}_{\text{SO}_4\text{-OH}}$ isotope fractionations, and homogenization temperatures given in Jannas et al. (1999). In general, the δD – $\delta^{18}\text{O}$ mineral data and calculated fluid compositions of main-stage fluids fall on trends (Figs. 6 and 7) nearly parallel to the meteoric water line. The trend is also similar to that observed for alunite in active stratovolcanoes (Zimbelman et al., this volume). These trends reflect mixing of magmatic and ^{18}O -enriched meteoric water and are in contrast to the typical mixing trends between magmatic and unexchanged meteoric water commonly observed for high-sulfidation ore-forming fluids (e.g., Julcani; Deen et al., 1994). To some extent, the range in δD values of alunite parental fluids may reflect the effect of significant deuterium enrichment of degassed aqueous fluid over silicate melts during Rayleigh degassing of different batches of magma throughout the period of hydrothermal alteration (Taylor, 1986, 1988, 1991). However, the data trends in Figs. 6 and 7 are not vertical, with a nearly constant $\delta^{18}\text{O}$, which may indicate that some mixing of unexchanged meteoric water with the magmatic fluids occurred in the different environments. Such mixing is most obvious in the shallow and late-stage fluids (Fig. 8).

6.1. Brecha Sylvestre, Stage 1, and Stage 2: magmatic-hydrothermal alunite

Fig. 6 summarizes the $\delta^{34}\text{S}$, δD , and $\delta^{18}\text{O}$ values for alunite from Brecha Sylvestre, Stage 1, and Stage 2.

Table 5
Sample descriptions and stable isotope results for all Tambo alunite, barite, and enargite samples

Sample	Lab no.	Stage	Deposit	Elev. (m)	Description	δD (‰)	$\delta^{34}S$ (‰)	$\delta^{18}O_{SO_4}$ (‰)	$\delta^{18}O_{OH}^a$ (‰)
<i>Alunite</i>									
WND-92A, 190.0 m	T09a	1	Wendy	4113	Qtz–alun alteration	–98	6.2	9.4	3.7
WND-92A, 290.8 m	T10a	1	Wendy	4135	Qtz–alun alteration	–53	22.1	15.8	10.1
KBD-85, 16.0 m	T16a	1	Kimberly	4113	Qtz–alun–pyl alt		7.2		
KBD-85, 306.8 m	T29a	1	Kimberly	3887	Qtz–alun alteration	–32	23.1	17.2	14.3
KBD-85, 94.1 m	T30a	1	Kimberly	4052	Qtz–alun alteration	–30	1.1	20.3	14.3
WND-92a, 82.7 m	T31a	1	Wendy	4090	Qtz–alun–dick alteration		12.4		
RA-01	T01a	2	Reina	4010	Vein alun with bar, walth, Au	–28	27.3	21.0	15.5
RA-4, 41.6 m	T24a	2	Reina	4025	Coarse alun with walth	–27	27.4	20.3	16.3
CS-29	T26a	2	Canto Sur	4522	Fine-grained alun, bar with Au	–55	27.3	16.5	10.8
CS-01b	T27a	3	Canto Sur	4474	Alun with bar in breccia matrix	–38	2.1	19.6	15.3
CS-08	T02b	3	Canto Sur	4474	Matrix alun over bar	–78	1.8	10.1	3.6
LN-24	T03a	3	Kimberly West	4130	Matrix alun, overgrowing T03b	–47	5.5	17.7	13.2
LN-24	T03b	3	Kimberly West	4130	Matrix alun, overgrowing qtz	–46	1.3	19.2	12.8
WND-86, 155.0 m	T04a	3	Wendy	4017	Matrix alun, with enarg and scord	–63	1.7	15.0	8.1
WND-86, 237m	T06a	3	Wendy	3960	Matrix alun hosting enarg	–85	1.7	11.9	5.7
CS-04	T20a	3	Canto Sur	4475	Matrix alun	–41	–0.2		
WND-92a, 37.5 m	T21a	3	Wendy	4080	Matrix alun hosting enarg	–80	1.7	15.9	4.9
WND-92a, 231.8 m	T22a	3	Wendy	4122	Matrix alun	–54	2.8	19.7	14.7
KB-43	T23a	3	Kimberly	4075	Matrix alun	–79	0.7		
LN-09	T25a	2	Kimberly West	4240	Matrix alun with bar, qtz	–42	25.7	19.6	16.4
WND-92a, 13.7 m	T28a	3	Wendy	4075	Fine-grained matrix alun	–98	2.8	11.1	3.6
KB-08	T36a	3	Kimberly	4050	Matrix alun with scord	–51	3.5	18.0	12.0
KB-47	T07a	MS	Kimberly		Coarse banded vein with minor hem	–40	0.5	20.5	14.0
KB-33B	T08a	MS	Kimberly	4100	Coarse banded vein with hem	–48	1.1	16.4	8.8
WN-07	T32a	MS	Wendy	4180	Banded vein with hem	–69	1.4	15.9	9.7
KB-01	T33a	MS	Kimberly	4050	Coarse banded vein	–60	0.4	18.8	15.6*
KB-44	T34a	MS	Kimberly	4075	Coarse banded vein with hem	–74	0.3	11.3	
KB-09	T14a	Ca	Kimberly	4060	Thin veinlet cross-cutting altered tuff	–118	1.8	11.7	3.2
LN-02	T35a	Ca	Kimberly West	4260	Thin veinlet cross-cutting altered tuff	–56	9.2	20.6	14.2
KB-37B	T13a	SH	Kimberly	4480	Near-surface qtz–kao–alun–S alteration	–53	5.9	18.8	12.3
KB-40	T17a	SH	Kimberly	4460	Near-surface qtz–kao–alun–S alteration	–13	4.9	23.2	18.8
WN-14	T18a	SH	Cerro Elephant	4450	Powdery qtz–alun veinlets	–62	16.6	15.4	11.9*
CS-08	T02a	LV	Canto Sur	4474	Fine-grained alun–jar over Stage 3 alun	–111	–0.2	10.0	2.5
KBD-85, 306.8 mV	T11a	LV	Kimberly	3887	Alun–jar veinlet in Stage 1 alteration	–110	5.4	7.0	1.3
KBD-85, 125.7 m	T12a	LV	Kimberly	4027	Alun–qtz (jar) veinlet	–57	2.0	15.7	9.8
WND-86, 226.5 m	T15a	LV	Wendy	3969	Brown jar–alun vein	–68	4.3	16.1	9.3
<i>Barite</i>									
RA-01	Tb01	2	Reina	4050	Coarse vein fill, associated with T01a		26.8	13.8	
CS-08	Tb02	2	Canto Sur	4474	Coarse matrix crystals, with alun		24.5	17.3	
CS-30	Tb03	2	Canto Sur	4396	Coarse matrix crystals, mineralized		23.8	17.1	
<i>Enargite</i>									
WND-86, 155.0 m	T04e	3	Wendy	4017	Enarg intergrown with alun	–2.4			
WND-86, 237 m	T06e	3	Wendy	3960	Enarg intergrown with alun	–3.7			

Elevation given in meters above sea level. Abbreviated mineral names: alunite (alun), barite (bar), dickite (dick), enargite (enarg), gold (Au), hematite (hem), jarosite (jar), kaolinite (kao), pyrophyllite (pyl), quartz (qtz), scorodite (scord), native sulfur (S), walthierite (wal).

^a Calculated using BrF₅ total oxygen data, except samples marked by *.

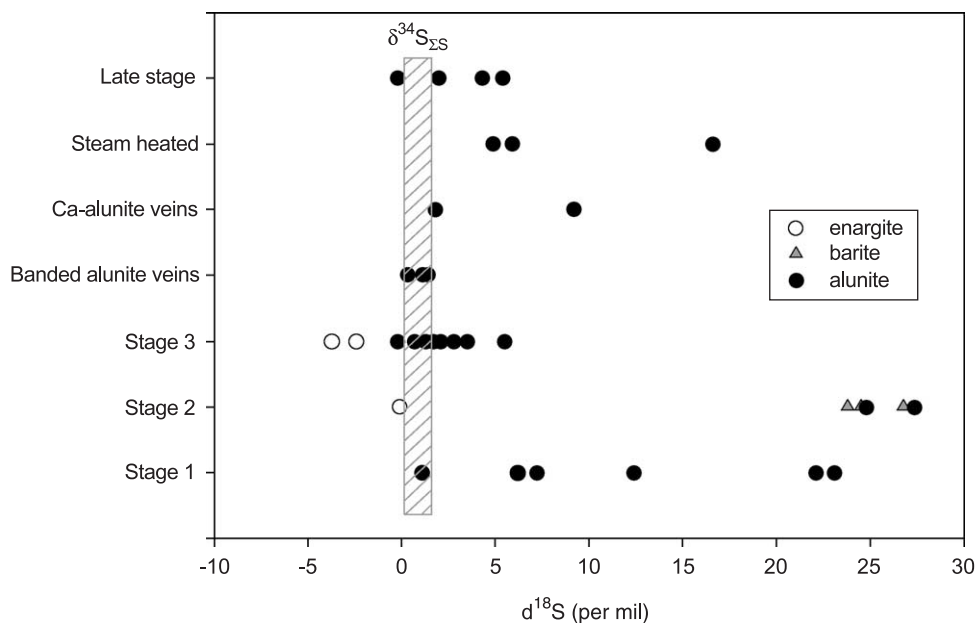


Fig. 5. Range in $\delta^{34}\text{S}$ data (in ‰) for all stages of Tambo alunite. Also shown are data for associated enargite (this study; Jannas et al., 1999) and barite from Stage 2 and Stage 3. Estimated $\delta^{34}\text{S}_{\Sigma\text{S}} = -1\%$ to 3% . See text for discussion.

Stage 1 $\delta^{34}\text{S}$ values range from 1% to 23% (Fig. 6A). Stage 2 alunite is heavier, with $\delta^{34}\text{S}$ values of 26% to 27% . Sulfur-isotope values for the two Brecha Sylvestre samples (14% and 15%) are lower than those for Stage 2 alunite, but are intermediate to those for Stage 1.

Values of δD and $\delta^{18}\text{O}$ for alunite are shown in Fig. 6B, along with the range of calculated fluid compositions in equilibrium with samples from each alteration event. The composition of both the Stage 2 and Brecha Sylvestre alunite parental fluids overlap those typical of fluids in felsic magmas (Taylor, 1988). Most Stage 1 fluids also plot within this range, indicating a dominant magmatic component. In both Stages 1 and 2, there is a linear trend to lower isotopic values as discussed above.

Discussion: High $\delta^{34}\text{S}$ values for Stage 2 alunite (and barite) and fluid compositions in the magmatic range are consistent with a magmatic-hydrothermal origin for this stage of alteration. In contrast, the large range in Stage 1 $\delta^{34}\text{S}$ values is unusual. The largest $\delta^{34}\text{S}$ values (22% to 23%) are consistent with derivation from sulfate that equilibrated with H_2S during disproportionation of SO_2 (Rye et al., 1992;

Rye, 1993), but other values are not. We propose two alternative explanations for this variability. The values may represent magmatic-hydrothermal alunite formed from aqueous sulfate-sulfur species that did not always reach isotopic equilibrium with H_2S during disproportionation of SO_2 . This could result from shorter residence times for aqueous sulfate during early vapor-plume development, condensation, and alunite precipitation in Stage 1. Alternatively, the values can be interpreted to indicate magmatic-hydrothermal alunite that is partly to completely replaced by either magmatic-steam or steam-heated alunite. On the basis of the textural and petrographic observations for Stage 1 alunite, the second model is preferred. Alunite with the lowest $\delta^{34}\text{S}$ values has macroscopic characteristics of Stage 1 alteration, but these grains appear to have precipitated in vugs created by the leaching of feldspar phenocrysts. Partial replacement of APS-bearing alunite by a more homogeneous, K-rich variety is also observed in one sample. Similar processes are inferred at El Salvador (Watanabe and Hedenquist, 2001), where partial replacement of hypogene by supergene alunite resulted in intermediate $\delta^{34}\text{S}$ values. One

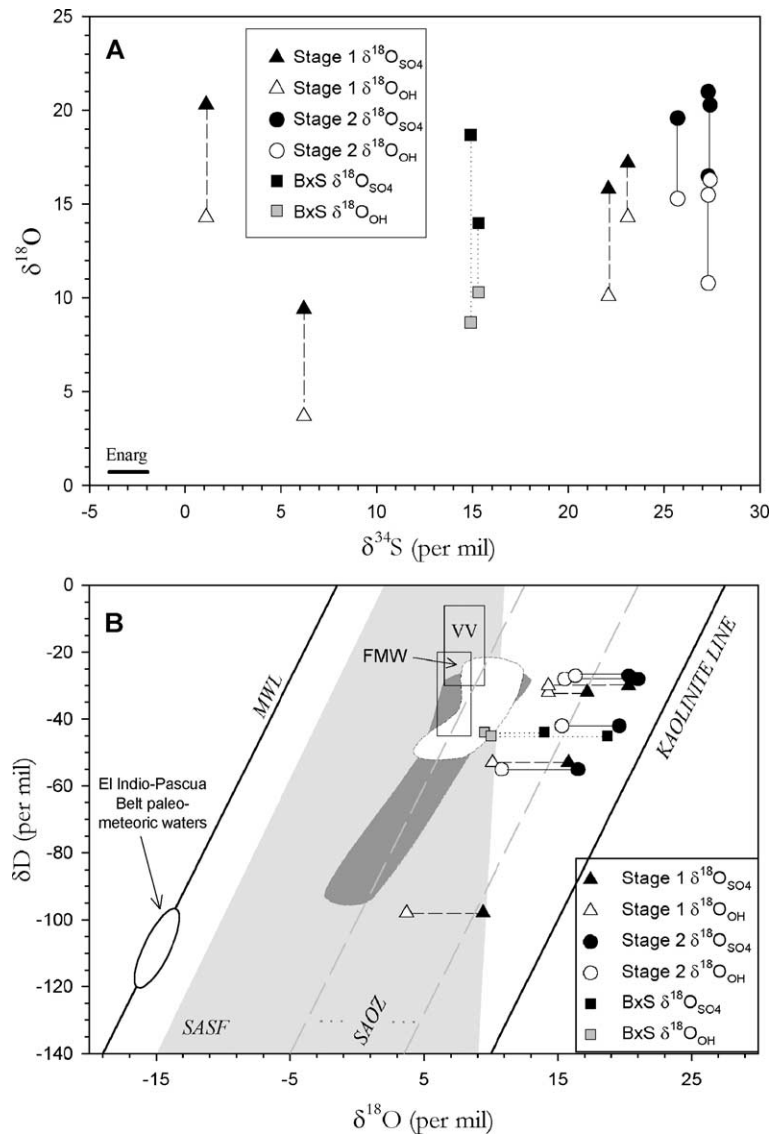


Fig. 6. (A) Data for $\delta^{34}\text{S}$ and $\delta^{18}\text{O}$ (in ‰) for alunitic from Stage 1, Stage 2, and Brecha Sylvestre (BxS). The $\delta^{18}\text{O}_{\text{SO}_4}$ and $\delta^{18}\text{O}_{\text{OH}}$ values for each sample are joined by lines. Range of $\delta^{34}\text{S}$ data for associated sulfides is also given. (B) Values of δD , $\delta^{18}\text{O}_{\text{SO}_4}$, and $\delta^{18}\text{O}_{\text{OH}}$ for alunitic from Stage 1, Stage 2, and Brecha Sylvestre (BxS). Ranges of calculated $\delta\text{D}_{\text{H}_2\text{O}}$ and $\delta^{18}\text{O}_{\text{H}_2\text{O}}$ for fluids in equilibrium with each stage are shown as shaded fields (gray=Stage 1, 200–280 °C; white=Stage 2, 180–250 °C). Lines and fields are: MWL=meteoric water line of Craig (1961); kaolinite line of Savin and Epstein (1970); V.V. (volcanic vapor)=range of water compositions discharged from high-temperature fumaroles (Giggenbach, 1992); SASF (light gray)=supergene alunitic SO_4 field; and SAOZ=supergene alunitic OH zone as described by Rye et al. (1992). The white circle is the estimated composition of paleo-meteoric waters.

alunitic sample has a significantly lower $\delta\text{D}_{\text{H}_2\text{O}}$ (−94‰) and larger $\delta^{18}\text{O}_{\text{SO}_4}$ values (7.0‰), which suggests mixing of isotopically heavy magmatic fluids with ^{18}O -enriched meteoric fluids (due to exchange, boiling, or evaporation).

6.2. Stage 3 and banded alunitic veins; transitional and magmatic-steam alunitic

Fig. 7 summarizes the $\delta^{34}\text{S}$, δD , and $\delta^{18}\text{O}$ values for Stage 3 and banded-vein alunitic. The $\delta^{34}\text{S}$ values

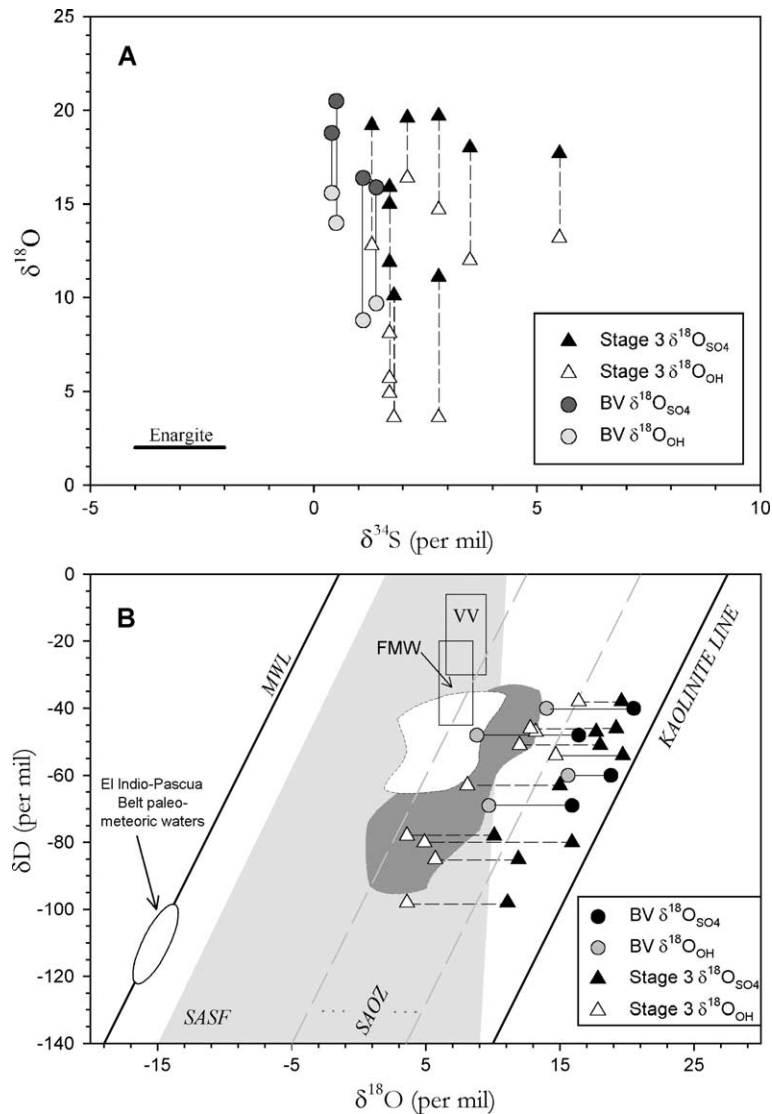


Fig. 7. (A) Data for $\delta^{34}\text{S}$ and $\delta^{18}\text{O}$ (in ‰) for Stage 3 and banded vein (BV) alunite. The $\delta^{18}\text{O}_{\text{SO}_4}$ and $\delta^{18}\text{O}_{\text{OH}}$ values for each sample are joined by lines. Range of $\delta^{34}\text{S}$ data for Tambo sulfides is also given. (B) Values of δD , $\delta^{18}\text{O}_{\text{SO}_4}$, and $\delta^{18}\text{O}_{\text{OH}}$ for Stage 3 and banded vein (BV) alunite. Range of calculated $\delta\text{D}_{\text{H}_2\text{O}}$ and $\delta^{18}\text{O}_{\text{H}_2\text{O}}$ for fluids in equilibrium with each stage is shown as shaded fields (gray=Stage 3, 200–300 °C; white=BV, 200–300 °C). Fluid compositions calculated from equations of Stoffregen et al. (1994). Lines and fields are as given in Fig. 6.

for both events are slightly greater than the values of Stage 3 enargite (about -2‰ to -4‰). Banded-vein alunite has consistently lower $\delta^{34}\text{S}$ values (about 0‰ to 1.5‰) than Stage 3 samples. Only minor variation in $\delta^{34}\text{S}$ is reported for different banded-vein samples or for growth bands in a single vein (Deyell, 2001). The narrow range of values for both banded-vein and Stage 3 alunite indicates that sulfur-isotope equi-

librium was not attained between aqueous sulfate and H_2S .

Calculated fluid compositions for alunite are shown in Fig. 7B. Stage 3 fluids have $\delta\text{D}_{\text{H}_2\text{O}}$ values between about -30‰ and -95‰ , and $\delta^{18}\text{O}_{\text{H}_2\text{O}}$ values between -1‰ and 13‰ . The values for fluids from banded-vein alunite have a more restricted range, with $\delta\text{D}_{\text{H}_2\text{O}}$ between -35‰ and -65‰ . Data

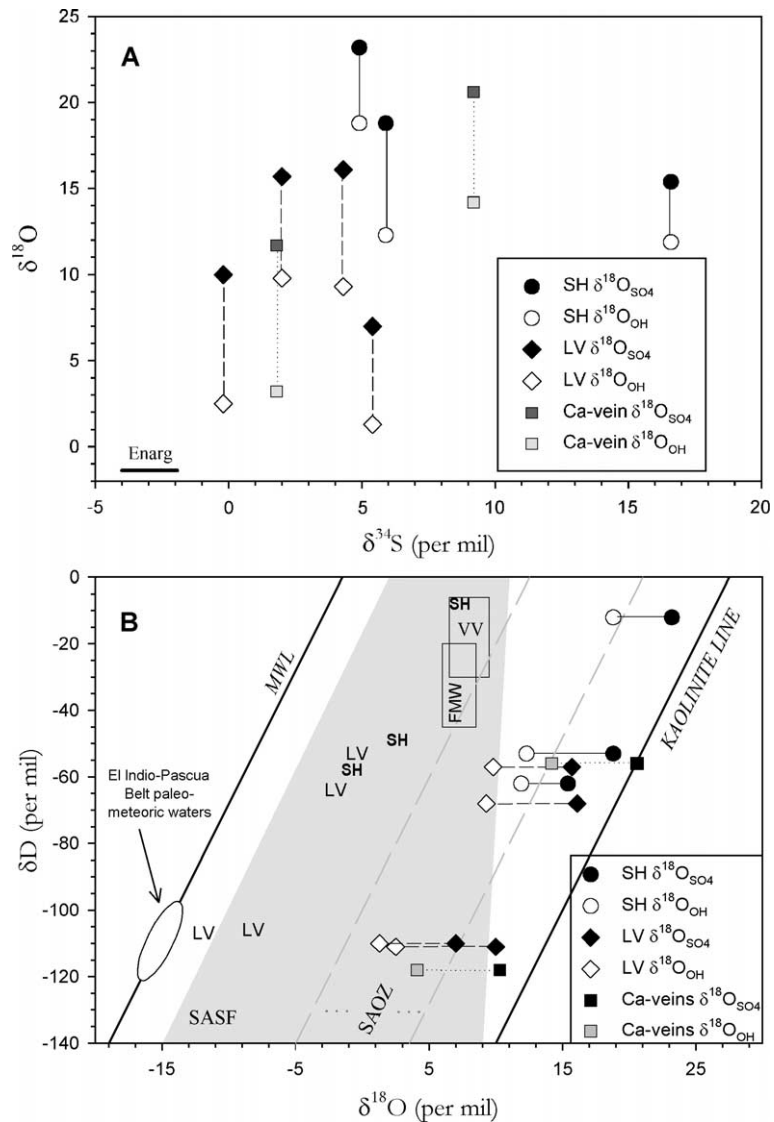


Fig. 8. (A) Data for $\delta^{34}\text{S}$ and $\delta^{18}\text{O}$ (in ‰) for steam-heated (SH), late vein (LV), and huangite-bearing vein (Ca-vein) alunite. The $\delta^{18}\text{O}_{\text{SO}_4}$ and $\delta^{18}\text{O}_{\text{OH}}$ values for each sample are joined by lines. Range of $\delta^{34}\text{S}$ data for Tambo sulfides is given for reference. (B) Values of δD , $\delta^{18}\text{O}_{\text{SO}_4}$, and $\delta^{18}\text{O}_{\text{OH}}$ for steam-heated (SH), late vein (LV) and huangite-bearing vein (Ca-vein) alunite. Ranges of calculated $\delta\text{D}_{\text{H}_2\text{O}}$ and $\delta^{18}\text{O}_{\text{H}_2\text{O}}$ for fluids in equilibrium with each stage are shown as symbols (SH=steam-heated fluids, 130 °C; LV=late-vein fluids, 80 °C). Lines and fields are as given in Fig. 6.

for both events overlap those typical of magmatic waters in felsic magmas (Taylor, 1988), but the fluids for Stage 3 and the banded veins are typically depleted in deuterium and enriched in ^{18}O relative to Stage 2 magmatic-hydrothermal fluids.

Discussion: Data for banded-vein alunite indicate a major magmatic component in the source

fluids. Results are consistent with a magmatic-steam origin. Stage 3 data are more variable and show greater degrees of mixing between magmatic and meteoric fluids. The larger range of $\delta^{34}\text{S}$ values for alunite of this stage suggests limited exchange between aqueous sulfur species. Deposition is thought to have occurred in an environment

that is transitional between magmatic-hydrothermal and magmatic-steam.

Depositional temperatures calculated from $\Delta^{18}\text{O}_{\text{SO}_4\text{-OH}}$ for banded-vein alunite average about 110 °C, with the exception of one higher value at 320 °C (sample T33a). Independent estimates of temperature are not available for this stage of alteration, but temperatures are expected to be at least as high as those encountered in the magmatic-hydrothermal environment (i.e., 180–300 °C). The effects related to retrograde isotopic exchange are not expected given the coarse grain size of the alunite; hence, the origin of the low values for $\delta^{18}\text{O}_{\text{OH}}$ is unclear. Similar behavior has been reported for magmatic-steam alunite at Alunite Ridge, Marysvale, UT (Rye et al., 1992).

The range in $\Delta^{18}\text{O}_{\text{SO}_4\text{-OH}}$ temperatures for Stage 3 alunite is greater than for the banded-veins (10–330 °C), but most values are below 200 °C. Again, several temperatures are unreasonably low but others are within the expected range for shallow environments of deposition.

As discussed above, the correlation between decreasing $\delta\text{D}_{\text{H}_2\text{O}}$ and $\delta^{18}\text{O}_{\text{H}_2\text{O}}$ values for both alunite types does not project back to the meteoric water line. This trend can be attributed to mixing of magmatic and ^{18}O -enriched (from exchange, boiling, or evaporation) meteoric water.

6.3. Huangite-bearing vein alunite

The $\delta^{34}\text{S}$, δD , and $\delta^{18}\text{O}$ values for two samples of huangite-bearing vein alunite are plotted in Fig. 8. Results for the two samples are significantly different. One sample has low $\delta^{34}\text{S}$ (1.8‰) and $\delta^{18}\text{O}_{\text{SO}_4}$ (11.7‰) values. The other sample has a $\delta^{34}\text{S}$ value of about 9‰, and has a $\delta^{18}\text{O}$ value in the range of magmatic-hydrothermal alunite. Calculated $\delta\text{D}_{\text{H}_2\text{O}}$ values in equilibrium with the two samples are –115‰ and –52‰, respectively. The first sample (T35) has a fluid composition which overlaps that of Tambo magmatic-hydrothermal and magmatic-steam fluids. Data for the latter (sample T14a) indicate a significant ^{18}O -enriched meteoric fluid component or a highly fractionated late-stage magmatic fluid with a minor component of unexchanged meteoric water.

Discussion: Depositional temperatures for the two huangite-bearing samples, on the basis of $\Delta^{18}\text{O}_{\text{SO}_4\text{-OH}}$

values, are similar and average 105 °C. No independent temperature estimates are available for these veins, however, and it is unclear whether these values are reasonable. Temperatures of formation for the mineralogically similar El Indio Campana vein were estimated to be 225–275 °C (Jannas, 1995), but lower temperature occurrences of Ca-rich alunite have been reported elsewhere (e.g., Stoffregen and Alpers, 1987; Arribas et al., 1995). The origin of these veins and their relation to other alteration or mineralizing events is uncertain.

6.4. Near-surface alteration

Fig. 8 summarizes the $\delta^{34}\text{S}$, δD , and $\delta^{18}\text{O}$ data for alunite sampled from the near-surface alteration blanket on Cerro Elefante. Two samples (T13, T17) have $\delta^{34}\text{S}$ values slightly enriched relative to those of the bulk sulfur in the deposit, and have high $\delta^{18}\text{O}_{\text{SO}_4}$ values. The third sample (T18), selected from white, powdery veins that crosscut Stage 1 alteration just below the blanket zone, has a much larger $\delta^{34}\text{S}$ value (16.6‰) and lower $\delta^{18}\text{O}_{\text{SO}_4}$ value (15.4‰) than those of the first two samples.

Calculated fluid compositions in equilibrium with alunite are plotted in Fig. 8B. Values of $\delta\text{D}_{\text{H}_2\text{O}}$ for the first two samples vary between about –10‰ and –55‰, with $\delta^{18}\text{O}_{\text{H}_2\text{O}}$ between 2‰ and 7‰. These values are significantly enriched relative to those for meteoric water, and indicate a dominant magmatic fluid component. Deuterium enrichment by evaporation (Henley and Stewart, 1983) is inferred for the highest δD value.

Discussion: Sulfur and oxygen isotopic data for this stage of alteration are consistent with a steam-heated origin. Limited sulfur-isotope exchange between aqueous SO_4 and H_2S is inferred for the first two samples. Calculated $\Delta^{18}\text{O}_{\text{SO}_4\text{-OH}}$ temperatures of deposition are 110 and 210 °C. Results are reasonable, although perhaps slightly high in the latter, for environments in which aqueous sulfate forms at shallow levels from the oxidation of H_2S by atmospheric oxygen (Rye et al., 1992; Ebert and Rye, 1997). Oxygen isotopic temperatures also suggest that oxygen-isotope equilibrium was obtained between the aqueous sulfate and water in the fluid. The third sample, given its shallow depth, has an unreasonably high $\Delta^{18}\text{O}_{\text{SO}_4\text{-OH}}$ temperature of 290 °C, although $\delta^{18}\text{O}_{\text{OH}}$ values are in the range of those for

the previous two samples. Significant isotopic exchange of S occurred between aqueous SO_4 and H_2S , but $\Delta^{18}\text{O}_{\text{SO}_4\text{-OH}}$ may have also been affected by post-depositional exchange.

6.5. Alunite in late veins

Sulfur data for late-vein alunite are slightly greater than values for sulfides in the deposit (Fig. 8A). All $\delta^{18}\text{O}_{\text{OH}}$ values plot within the Supergene Alunite OH Zone (SAOZ) defined by Rye et al. (1992); however, all but one of the $\delta^{18}\text{O}_{\text{SO}_4}$ values plot outside of the Supergene Alunite Sulfate Field (SASF; Rye et al., 1992). These results, combined with large positive $\Delta^{18}\text{O}_{\text{SO}_4\text{-OH}}$ values, are not consistent with a supergene origin. Similarly, calculated $\Delta^{18}\text{O}_{\text{SO}_4\text{-OH}}$ temperatures average about 100 °C and are higher than expected for deposition in the supergene environment.

Calculated fluid compositions in equilibrium with alunite samples are shown in Fig. 8B. The data fall on a trend that projects, at low δD values, close to the meteoric water line. Two samples with the lowest $\delta\text{D}_{\text{H}_2\text{O}}$ and $\delta^{18}\text{O}_{\text{H}_2\text{O}}$ values have fluids close to the predicted composition of meteoric waters. The other two samples have much larger $\delta\text{D}_{\text{H}_2\text{O}}$ and $\delta^{18}\text{O}_{\text{H}_2\text{O}}$ values, indicating a substantial magmatic contribution. These fluid compositions overlap those for magmatic-hydrothermal and magmatic-steam occurrences.

Discussion: Results indicate that the late alunite veins were not generated under typical supergene conditions even though they have a field appearance suggestive of a supergene origin. This type of alunite formed from mixed magmatic–meteoric fluids at moderate temperatures, ~100 °C. The meteoric water component was only moderately enriched in ^{18}O . The small amount of sulfides in the orebody and surrounding host rock may indicate that some of the aqueous sulfate for the late alunite was derived from oxidation of H_2S .

7. Geochemistry of alunite

Major- and trace-element analyses were determined to evaluate the compositional variability of alunite and alunite-group minerals in each alteration stage. End-

member alunite is $\text{KAl}_3(\text{SO}_4)_2(\text{OH})_6$, but naturally occurring minerals of the alunite supergroup can have a wide range of chemical substitutions (Jambor, 1999). The most common substitution of Na for K defines the alunite–natroalunite solid solution. Other substitutions for K^+ may include Ca, Ba, Sr, REE, Pb, Ag, H_3O , and NH_4 . Substitution of Fe^{3+} for Al^{3+} defines the alunite–jarosite solid solution, and substitution of $(\text{PO}_4)^{3-}$ for $(\text{SO}_4)^{2-}$ forms APS minerals of the crandallite and beudantite groups (Table 3: as summarized in Jambor, 1999). Several experimental (Hemley et al., 1969; Stoffregen and Cygan, 1990) and empirical studies (e.g., Aoki, 1991; Aoki et al., 1993; Thompson, 1992; Hedenquist et al., 1994; Arribas et al., 1995) have correlated variations in alunite-group chemistry to specific environments or temperatures of deposition. Similarly, published data on REE contents of acid-sulfate-altered rocks suggest that REE contents may be influenced by a number of factors, including (a) crystallographic controls (Schwab et al., 1990); (b) the availability and type of complexing agents (Wood, 1990a,b; Lottermoser, 1992); and (c) environmental controls such as pH, Eh, and temperature (e.g., Michard, 1989). REE partitioning between magmatic vapors and melts (Flynn and Burnham, 1978; Candela and Piccoli, 1995; Reed, 1995) may also affect the REE content of any alteration or ore minerals that form from the condensation of magmatic fluids in the epithermal environment.

7.1. Summary of results

Chemical data for each stage of Tambo alunite are given in Tables 6a and 6b, which includes a summary of EPMA data for all analyses of alunite, and ICP-MS trace-metal concentrations for selected samples. REE results in this study (Fig. 9) are normalized to the average REE contents of unaltered host rock (Tilito Formation: Table 2). Our interpretation of EPMA results is limited by the large beam size required for analysis (see Section 3). Most fine-grained alunite samples, particularly supergene and steam-heated samples, were too small to be analyzed by this method and relatively few good analyses were obtained. Similarly, thin oscillatory zones and growth bands (locally <10 μm) could not be analyzed. In such cases, results represent an average over multiple compositional bands.

Table 6a
Summary of EPMA data for all Tambo alunite-group samples

	Stage 1				Stage 2				Stage 3			
	<i>n</i> =85 (5 samples)				<i>n</i> =59 (4 samples)				<i>n</i> =76 (3 samples)			
	Mean	2 σ	Max	Min	Mean	2 σ	Max	Min	Mean	2 σ	Max	Min
K	0.81	0.46	1.03	–	0.58	0.80	1.04	0.02	0.89	0.31	1.05	0.26
Na	0.19	0.34	0.81	–	0.09	0.27	0.90	–	0.11	0.26	0.62	–
Ca	0.03	0.27	0.83	–	–	–	0.02	–	–	–	0.02	–
Ba	–	–	0.03	–	0.18	0.40	0.51	–	–	–	0.06	–
Sr	–	–	0.09	–	–	–	–	–	–	–	–	–
Fe	–	–	0.06	–	0.04	0.55	2.13	–	–	–	0.17	–
Al	3.08	0.12	3.25	2.95	3.11	0.55	3.26	1.14	3.16	0.13	3.29	3.01
P	0.05	0.43	1.35	–	–	–	0.05	–	–	–	0.06	–
S	1.95	–	2.00	0.65	1.99	0.02	2.00	1.95	1.99	0.03	2.00	1.94
	Banded vein				Steam-heated				Late-stage			
	<i>n</i> =33 (2 samples)				<i>n</i> =31 (2 samples)				<i>n</i> =11 (2 samples)			
	Mean	2 σ	Max	Min	Mean	2 σ	Max	Min	Mean	2 σ	Max	Min
K	0.87	0.20	1.02	0.61	0.98	0.10	1.04	0.84	0.92	0.10	1.02	0.85
Na	0.10	0.15	0.33	–	0.04	0.08	0.16	–	0.12	0.10	0.21	0.02
Ca	–	–	–	–	–	–	0.03	–	–	–	0.02	–
Ba	–	–	0.04	–	–	–	–	–	–	–	–	–
Sr	–	–	0.02	–	–	–	–	–	–	–	–	–
Fe	–	–	–	–	–	–	0.02	–	0.04	0.11	0.21	–
Al	3.14	0.11	3.24	3.04	3.13	0.09	3.23	3.05	3.13	0.18	3.25	2.94
P	0.02	0.04	0.08	–	–	–	0.03	–	–	–	0.04	–
S	1.98	0.04	2.00	1.92	1.99	0.02	2.00	1.97	1.99	0.02	2.00	1.96

Results are in atoms per formula unit (a.p.f.u.) calculated for (P,S)O₄=2 (Jambor, 1999). Data below detection limits (0.02 a.p.f.u. for Na and 0.01 for all other elements) are not shown. *n*=total number of analyses.

Magmatic-hydrothermal alunite, which includes Brecha Sylvestre, Stage 1, and Stage 2 samples, exhibits a wide range of compositions. Samples of Brecha Sylvestre alunite are compositionally simple, with local enrichment in Pb and Sb, minor Na, and trace P, Sr, and Ba. Alunite is depleted in REE relative to host-rock values, and moderate fractionation in heavy-REE (HREE) is apparent. In contrast, Stage 1 alunite is chemically heterogeneous, with irregular trace-element (Na, Ca, Ba, P, Sr) substitution. Average Na contents are the highest among the analyzed alunite types (statistically significant at 95% confidence limit, based on *t*-test results). Substitutions by Ca and Ba locally approach huangite and walthierite end-member compositions. REE results show Stage 1 alunite samples are depleted in mid-REE (MREE) relative to the host Tilito Formation, and exhibit minor light-REE (LREE) fractionation.

Stage 2 alunite has a unique geochemical signature characterized by high Ba and the presence of walthierite. Compositions intermediate between walthierite and alunite are common, and a nearly complete range of solid-solution compositions was measured (Fig. 10). Stage 2 alunite in the Kimberly and Canto Sur deposits is commonly oscillatory zoned (Fig. 4c), typically with a core of Na-bearing alunite overgrown by walthierite and rimmed by nearly stoichiometric alunite. Walthierite has not been identified in the Wendy breccia, and Stage 2 alunite in this area occurs as coarse bladed crystals that are zoned in Ca, Na, P, and minor Ba. Stage 2 alunite is typically Na-poor (<0.2 atoms per formula unit—a.p.f.u.), although the mineral locally approaches natroalunite end-member compositions. Barium, Pb, Sb, and Ag contents on average are the highest in Stage 2 compared to the contents in other alunite types. Stage 2 alunite

Table 6b
ICP-MS trace-element and REE concentrations (ppm) for bulk (~1 g) alunite samples

Sample	Stage 1 ^a		Stage 2		Stage 3			
	T09	T10	T01	T27	T03b	T04	T22	T28
P	970	380	1570	120	1180	1210	1700	1270
Sr	668	208	511	461	411	696	728	1580
Ba	340	630	4080	3850	3600	600	280	1470
Pb	116	856	1515	1745	251	608	428	338
Sb	–	11	317	10	10	29.5	31	15
Ag	2.7	0.9	79.4	4.0	5.8	21.4	0.2	3.0
Cu	1218	10	32	20	17	53	18	69
Bi	22	1.5	6.5	0.8	13.4	102.0	2.3	32.0
Zn	4	2	–	2	6	48	–	46
La	45.3	25.5	10.5	13.3	–	66.5	90.3	–
Ce	86.5	45	9.5	9.2	–	79.5	81.3	–
Pr	9.2	4.3	0.6	0.5	–	4.1	5.1	–
Nd	24.8	11.5	1.5	0.8	–	6	6.8	–
Sm	3.3	1.6	0.2	–	–	0.2	0.1	–
Eu	0.3	0.3	0.2	0.2	–	0.2	0.1	–
Gd	1.8	1.3	0.1	–	–	0.3	0.1	–
Tb	0.2	0.3	–	–	–	–	–	–
Dy	1	1.1	–	–	–	0.2	–	–
Ho	0.2	0.3	–	–	–	–	–	–
Er	1	1.1	–	–	–	–	–	–
Tm	0.1	0.3	–	–	–	–	–	–
Yb	1.4	1.6	–	–	–	–	–	–
Lu	0.2	0.3	–	–	–	–	–	–

Sample	Bx Sylvestre	Banded vein		Ca-V	SH	Late-stage	
	B03	T33	T32	T14	T17	T11	T02A
P	680	2300	1410	630	940	1490	1580
Sr	511	1535	582	246	353	2760	677
Ba	650	1120	450	830	1030	2140	2430
Pb	1700	580	628	59	916	486	2240
Sb	78.5	28.5	22	0.5	10	14.5	14.3
Ag	0.8	–	0.6	–	1.4	3.0	7.9
Cu	25	18	19	18	44	44	1
Bi	29.2	0.3	46.2	0.6	1.0	32.3	6.0
Zn	2	–	2	8	18	70	–
La	33	156	81.5	48	47.5	29	77
Ce	3	198	116.5	109.5	85	50	47
Pr	8	13.4	7.9	15.5	8.7	5.9	2.1
Nd	0.6	24	13	65.5	27.5	22	3
Sm	0.1	1.1	0.6	16.8	3.7	5	0.2
Eu	0.5	0.3	0.1	2.5	0.5	0.9	0.4
Gd	–	1	0.5	9.3	2.3	3.2	0.1
Tb	0.1	0.1	–	0.6	0.2	0.4	–
Dy	–	0.3	0.1	0.7	0.7	1.1	–
Ho	0.1	–	–	–	0.1	0.1	–
Er	–	0.1	–	–	0.5	0.3	–
Tm	0.1	–	–	–	–	–	–
Yb	–	0.1	–	0.1	0.8	0.3	0.1
Lu	–	–	–	–	0.1	–	–

^a Trace contaminants may be present. Detection limits (ppm) are: P (10), Sr (0.2), Ba (0.5), Pb (2), Sb (2), Ag (0.05), Cu (0.02), Bi (0.01), Zn (2), REE (0.1). Data below detection not shown. Precision estimates, based on duplicate analyses, are better than stated value at 95% confidence limit, 90th percentile: La, Ce, Pr (5%), all other REE (10%), P, Sr, Sb, Ni (10%), Ba (15%), Ag, Cu (22%), Zn (55%).

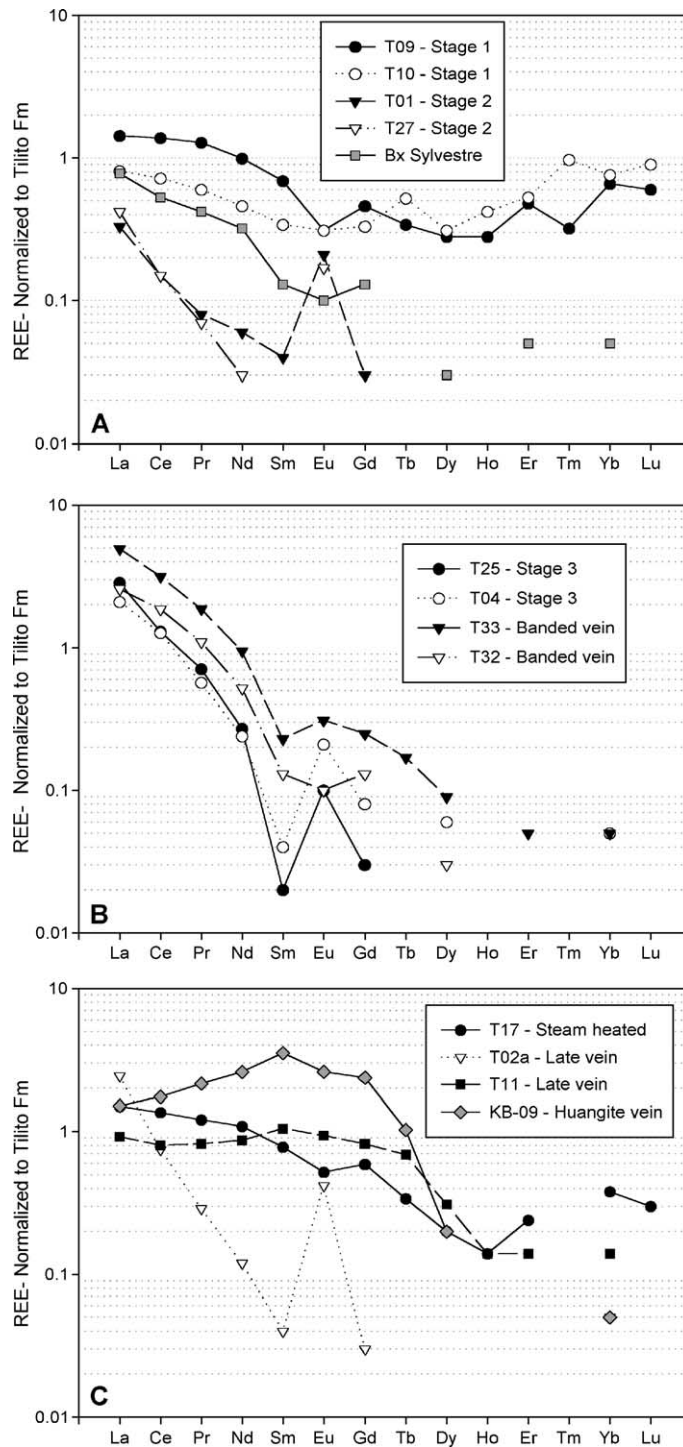


Fig. 9. REE data for all stages of Tambo alunite normalized to the average REE composition of the Tilito Formation. Individual sample numbers are given in the legend accompanying each plot. Data points below detection are not plotted. (A) Bx Sylvestre, Stage 1 and Stage 2 alunite. (B) Stage 3 and banded vein alunite. (C) Huangite-bearing vein alunite (Ca vein), steam-heated, and late-vein samples.

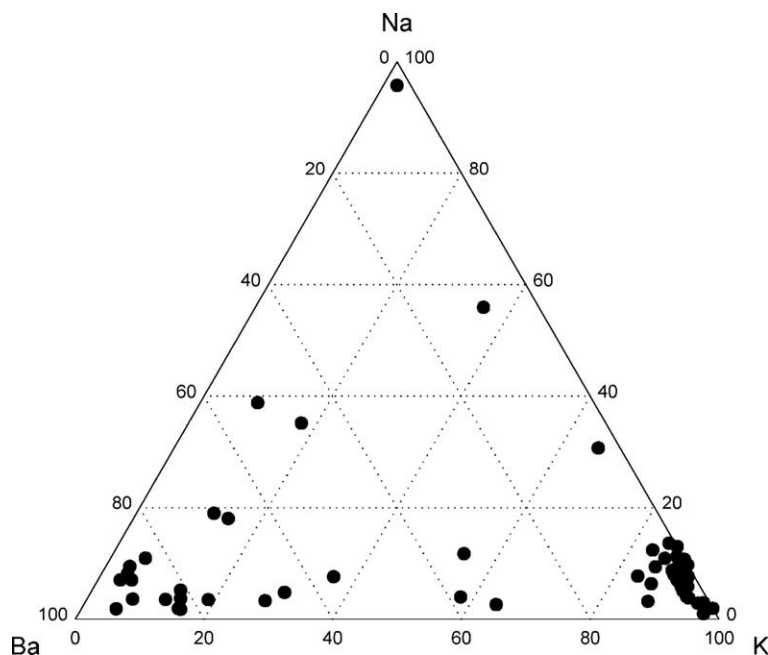


Fig. 10. Normalized K–Na–Ba compositions of Stage 2 alunite. Data points are recalculated EPMA compositions (58 analyses total from four samples).

exhibits strong fractionation in LREE, no detectable HREE, and a strong, positive Eu anomaly.

Compositions of Stage 3 and banded-vein alunite are nearly stoichiometric, with minor Na (<0.4 a.p.f.u) and trace-metal substitution. Oscillatory zoning with trace $\text{PO}_4 \pm \text{Sr}$ substitution is common, and banded-vein alunite in particular has the highest P contents compared with those in other stages of alunite. In contrast, Ca, Ba, and Pb contents are low. EPMA analyses indicate almost no chemical variation across successive growth bands in the alunite veins. Stage 3 and banded-vein alunite are both enriched in LREE, with values two to five times higher than those of the host rock, but are significantly depleted in HREE. These trends are more pronounced in the banded-vein samples with $(\text{La}/\text{Yb})_n$ ratios >100.

Chemical data for samples of steam-heated alunite are limited because of the small grain size and relatively rare occurrence of alunite in the steam-heated alteration zone. This alunite is nearly stoichiometric, with minor Na, Sr, and P substitution. Fractionation of HREE is moderate, with minor enrichment in LREE compared with that of the host Tilito Formation. Huangite in huanigite-bearing veins typically occurs as thick over-

growths on alunite grains that have woodhouseite–svanbergite cores. EPMA data indicate a full range of substitution between alunite and huanigite end-member compositions (Fig. 11). Oscillatory zoning between huanigite and minamiite compositions occurs locally. ICP-MS results confirm very high Ca contents, with elevated Na concentrations and few trace elements. Alunite of this stage is significantly enriched in LREE and MREE relative to all other alunite types. HREE are strongly fractionated.

Geochemical data for the late veins are limited because of the fine-grained nature of the alunite. Results indicate that the mineral is chemically variable, with high Sr contents and locally elevated Ba, Pb, Cu, Zn, and P. Sodium contents are low, and REE contents are extremely variable. One sample (T11) shows no LREE fractionation relative to host-rock values but has a significant depletion in HREE. Another sample (T02a) has a compositional variation similar to that of Stage 2 alunite, with extreme HREE fractionation and a large, positive Eu anomaly. EPMA analyses of extremely fine-grained, pseudocubic alunite from irregular veinlets indicate nearly stoichiometric compositions, minor Na substitution (<0.2

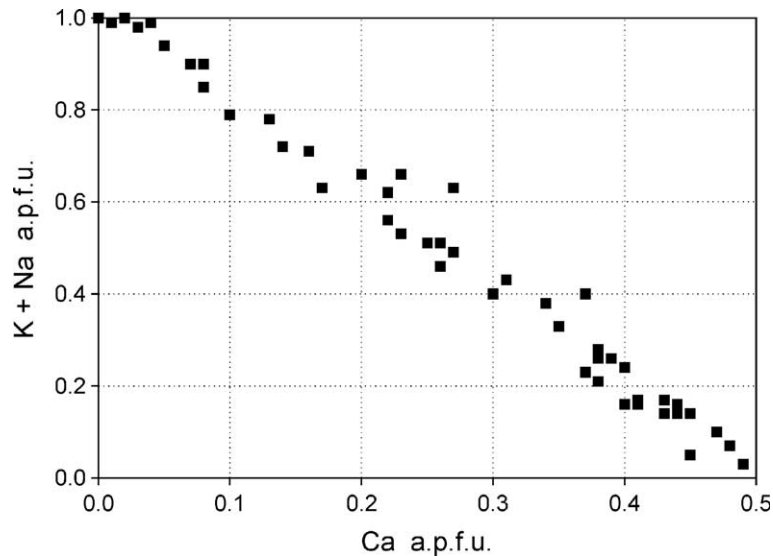


Fig. 11. Measured K+Na versus Ca content of huangite-bearing vein alunite. Axes are given as atoms per formula unit (a.p.f.u.). Data taken from EPMA analyses ($n=47$) of multiple grains in two samples, KB-09 and LN-02.

a.p.f.u.), and trace Fe. Slightly larger (10–15 μm) pseudocubic grains of Ca- and Sr-bearing APS minerals (likely woodhouseite–svanbergite) occur intergrown with alunite and kaolinite in these veins.

7.2. Interpretation

Chemical data for Tambo magmatic-hydrothermal alunite are consistent with other reports of heterogeneous alunite compositions and elevated concentrations of Na, Ba, Ca, Sr, PO_4 , and REE in high-temperature environments (Stoffregen and Alpers, 1987; Stoffregen and Cygan, 1990; Aoki, 1991; Aoki et al., 1993; Hedenquist et al., 1994; Thompson, unpublished data). These elements typically occur as APS minerals overgrown by alunite or alunite–natroalunite (e.g., Stoffregen and Alpers, 1987; Aoki et al., 1993), as has been observed in Stage 1 samples from this study.

Few chemical data are available for alunite from other near-surface occurrences. Limited data for steam-heated alteration show significant variations (Ebert and Rye, 1997; Thompson, unpublished data), with compositions extending to the alunite and natroalunite end-members. The former is consistent with results reported in this study. Supergene alunite is generally K-rich in compar-

ison with that in higher temperature occurrences, although trace-element concentrations are extremely variable, with Ca, Sr, REE, and PO_4 substitutions reported (Stoffregen and Alpers, 1987; Arribas et al., 1995). The elevated trace-metal concentrations for samples of late-vein alunite examined in this study are attributed to the metal content of source fluids because stable-isotope results are not consistent with the oxidation of precursor sulfides.

Few chemical analyses have been published for magmatic-steam alunite. Alunite in the Tambo banded veins, although chemically similar to that from Alunite Ridge, Utah, lacks the same degree of fine-scale, sawtooth bands of PO_4 and Sr substitution (Cunningham et al., 1996). Substitution of Ba is also more common in Alunite Ridge samples, and is rarely detected in the Tambo banded-vein alunite.

Recent studies of the REE contents of hydrothermal fluids (Michard, 1989; Lewis et al., 1997, 1998) and of hydrothermally altered rocks (Hopf, 1993; Arribas et al., 1995; Fulignati et al., 1999) have shown that REE can be mobilized under the conditions of acid-sulfate alteration. Compared with fresh-rock equivalents, magmatic-hydrothermal alunite has been characterized by strong fractionation of HREE with respect to LREE (Arribas et al., 1995; Fulignati et al., 1999). However, compositional data for REE in the host rocks and in

Stage 1 alunite at Tambo are not consistent with these published trends. We propose that the presence of APS inclusions and growth zones with elevated and Sr and P contents contribute to the ‘capturing effect’ described by Schwab et al. (1990) and allow for the incorporation of HREE into the alunite crystal structure. Similar behavior is noted for Tambo steam-heated alunite, which shows relatively minor HREE fractionation compared to host-rock values. Results are analogous to those described by Lewis et al. (1998) for altered rocks at Yellowstone National Park.

All alunite sampled from Stages 2 and 3 and the banded veins exhibits strong fractionation of HREE compared with the host-rock composition. This trend is most pronounced in the Stage 3 and banded-vein samples, whereas Stage 2 samples have comparatively lower REE concentrations. In all three stages, alunite was precipitated in open spaces (either in veins or breccia matrices), with little evidence for wallrock interaction. The chemistry of these alunite types should therefore reflect the composition of their source fluids. Experimental studies of REE partitioning between a Cl-bearing supercritical fluid and a residual melt have shown a relative enrichment in LREE compared with HREE in the vapor phase (Flynn and Burnham, 1978; Reed, 1995). No information is available for these phases at lower temperatures, but the chemical signature of the Stage 2, Stage 3, and banded-vein alunite may reflect the composition of a HREE-depleted fluid derived from the supercritical vapor-phase. In contrast, other forms of alunite derived their REE contents from a combination of magmatic fluids and REE leached from the host rocks.

8. Chemistry of fluid-inclusion gases

Bulk-sample fluid-inclusion techniques are widely used for the analysis of inclusion volatiles, mainly H₂O, CO₂, CH₄, N₂, H₂, and the noble gases (Shepherd and Rankin, 1998). Quadrupole mass spectrometric (QMS) analysis in particular has been applied to the identification of a large number of gases at low concentration in a variety of ore deposits and geothermal systems (e.g., Guha et al., 1990; Landis and Hofstra, 1991; Graney and Kesler, 1995b; Lindaas et al., 1998). A thorough review of this

technique and its limitations is given by Graney and Kesler (1995a).

Compositions of fluid-inclusion gas in Tambo alunite were determined by QMS analysis (Landis and Rye, *this volume*) to characterize Stage 2 and 3 ore fluids and banded-vein alunite parental (i.e., magmatic-steam) fluids. Repeated attempts to analyze fine-grained Stage 1, steam-heated, and late-vein alunite failed to release sufficient quantities of gas. We present herein only a summary of results, and the reader is referred to Landis and Rye (*this volume*) for details.

Alunite samples were selected from coarse-grained assemblages and were examined petrographically prior to analysis. Stage 2 alunite, selected from the Reina vein, contains mostly small (<10 μm), apparently primary, vapor-rich inclusions, but the results of necking are common. Other, slightly larger inclusions (10–15 μm) have irregular shapes and may be secondary or pseudo-secondary in origin. These inclusions are vapor-dominant, but with more variable liquid-to-vapor ratios. Stage 3 alunite was selected from the Canto Sur deposit and contains many small (5–10 μm), vapor-dominant inclusions. Banded-vein alunite, sampled from the Kimberly deposit, typically contains small, irregular, vapor-rich inclusions tightly clustered along growth zones. Repeated attempts to obtain fluid-inclusion homogenization temperatures for all three of these alunite stages were unsuccessful because of sample decrepitation.

8.1. Results and interpretation

Data for crush- and thermally-released gas from alunite samples are reported in Table 7. The quantity of gas released by crushing is significantly smaller than that by thermal release, and is likely the result of opening the very few and mostly larger inclusions that occur along secondary planes in the alunite. The genetic significance of these mechanically released gases is not known, but the gases probably represent mixtures of multiple pseudo-secondary generations and secondary trapping events. Thermally released gases are derived from decrepitation of the much more uniform occurrence of small primary inclusions (after removal of anomalous and secondary inclusions by crushing). These gases are generally considered to be

Table 7
Fluid-inclusion gas chemistry data for Stage 2, Stage 3, and banded vein alunite

Description	Stage 2		Stage 3		Banded Vein	
	Crush	Thermo	Crush	Thermo	Crush	Thermo
Weight (g)	8	8	6.6	6.6	7.4	7.4
N ₂ (%)	13.6131	0.1961	19.5394	0.5174	48.6233	20.2117
O ₂ (ppm)	48	44	19	59	127	1045
Ar (ppm)	4070	200	4431	654	14228	54992
H ₂ (%)	65.09	1.44	65.65	6.68	13.59	45.38
H ₂ O (%)	11.3008	97.551	4.0415	91.1306	21.7247	29.9172
H ₂ S (ppm)	20	17	12	12	22	1
SO ₂ (ppm)	38	3	24	30	43	48
HCl (ppm)	47	8	9	41	10	3
HF (ppm)	313	2672	90	2234	0	673
CH ₄ (%)	0.212	0.014	0.43	0.083	0.143	0.5
CO ₂ (%)	1.637	0.458	2.173	0.764	1.141	3.264
CO (%)	7.69	0.045	7.712	0.52	20.128	–
He (ppm)						44
⁴ He (V)	0.085718	1.479663	0.029593	0.945581	0.425582	0.107442
⁴ He (mol g ⁻¹)	6.872e-14	1.186e-12	2.876e-14	9.189e-13	3.689e-13	9.313e-14
³ He (cps)	2.771531	8.834217	1.057146	9.002538	17.72132	2.350376
³ He (mol g ⁻¹)	3.447e-20	1.099e-19	1.594e-20	1.357e-19	2.383e-19	3.161e-20
R= ³ He/ ⁴ He	5.016e-07	9.263e-08	5.542e-07	1.477e-07	6.460e-07	3.394e-07
R/R _a	0.3625	0.0669	0.4004	0.1067	0.4668	0.2452
²⁰ Ne	91.772	90.009	91.35	91.564	85.109	90.098
²¹ Ne	0.29924	0.28346	0.2571	0.25246	4.23851	0.39388
²² Ne	7.92907	9.70787	8.39272	8.1838	10.65245	9.50805
²¹ Ne/ ²⁰ Ne	0.0033	0.0031	0.0028	0.0028	0.0498	0.0044
²¹ Ne/ ²² Ne	0.0377	0.0292	0.0306	0.0308	0.3979	0.0414
²² Ne/ ²⁰ Ne	0.0864	0.1079	0.0919	0.0894	0.1252	0.1055

MH=magmatic-hydrothermal, MS=magmatic-steam, a=atmosphere, cps=counts per second. Blank fields: not detected.

representative of magmatic volatiles at the time of deposition (Graney and Kesler, 1995a). However, because of the analytical effects, the measured H₂S concentrations are considered to be a minimum value (Graney and Kesler, 1995b).

Ratios of major gas species, H₂S–SO₂–HCl (Fig. 12a), for the three alunite depositional events illustrate a distinct transition in volatiles chemistry. Stage 2 fluids are H₂S-dominant (H₂S/SO₂≈6), whereas magmatic-steam alunite (banded-vein sample) is characterized by a disequilibrium gas assemblage with high H₂, abundant reduced carbon species, and high SO₂. This chemical shift is also illustrated in Fig. 12b, with compositions recast to modeled equilibrium mole fractions at approximated temperatures and pressures for each alunite stage (using the NASA-CEA program and thermochemical data set; Gordon and McBride, 1994; McBride and Gordon, 1996; for details, see Landis and Rye, this volume).

Helium isotopes can be used as a tracer in studies of fluid flow in hydrothermal systems because isotopic compositions of He from atmospheric, upper-mantle, and crustal or radiogenic sources differ substantially. The atmospheric ³He/⁴He ratio (R_a) is 1.384±0.01×10⁻⁶ (Mamyrin et al., 1969; Ozima and Podosek, 2002) and is typically used as a reference for comparison to ³He/⁴He (R) gas data in the form R/R_a (such that R/R_a=1 for atmospheric He). Mantle-derived He has R/R_a values greater than 8 (Craig and Lupton, 1981; Lupton, 1983) whereas crustal ⁴He is enriched from both radiogenic U–Th decay and nuclear spallation reactions, creating low R/R_a values of <0.01 in upper crustal rocks that have elevated contents of U–Th (Gerling et al., 1971; Morrison and Pine, 1955). Thermal-release ³He/⁴He and R/R_a in this study range from 9.3e⁻⁸ to 3.4e⁻⁷ and 0.06 to 0.25, respectively, and clearly represent intermediate mixed values both from crustal sources and from lower crust to mantle sources with higher ³He. The unique

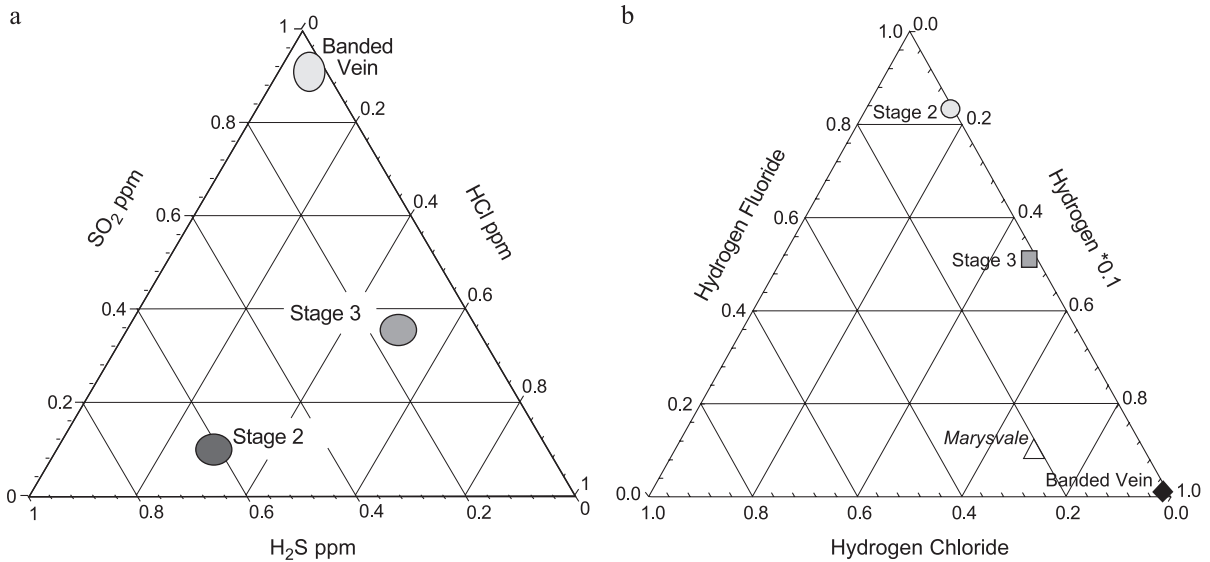


Fig. 12. (a) Ternary H₂S–SO₂–HCl data from fluid-inclusion analyses of Tambo alunite. (b) Ternary HCl–HF–H₂ data from fluid-inclusion analyses of Tambo alunite. Methods of recalculation of mole fractions are detailed in Landis and Rye (this volume).

chemistry of the gas in these alunite samples does not allow any involvement of atmospheric gas. Stable-isotope analyses indicate that only a minor amount of meteoric fluid was involved during any of these stages; hence, the volatiles and magmatic steam in particular must have been derived directly from magma, with He incorporated in the melt during partial melting in the lower crust or derived from condensed magmatic fluids that exchanged with crustal rocks. In particular, samples of magmatic-steam veins have the highest R/R_a values, the significance of which is discussed by Landis and Rye (this volume). We consider the highly disequilibrium gas chemistry with low He to be evidence that magmatic-steam alunite was formed from vapors that were boiled off from condensed magmatic fluids whose original gases had already been partly lost.

Similarly, the isotopic and elemental composition of Earth's Ne was different from that of the present atmosphere, and the difference can provide further indications of fluid source and the amount of crustal versus magmatic interaction. Neon isotopes are typically compared to those of end-member compositions of the atmosphere and to those of mid-ocean-ridge basalts (MORB), which show enrichment in ²⁰Ne and ²¹Ne relative to ²²Ne (e.g., Poreda and Radicati di Brozolo, 1984; Marty, 1989). In contrast,

crustal fluids typically are enriched in ²¹Ne and ²²Ne relative to ²⁰Ne from crustal nucleogenic–radiogenic processes (Kennedy et al., 1990). Neon isotopic data from this study are consistent with He results and suggest a crustal enrichment in ²¹Ne and ²²Ne relative to ²⁰Ne (Table 7).

9. Discussion

Fluid-inclusion and isotopic studies of several high-sulfidation deposits have indicated that magmatic fluids contribute to both alteration and mineralization within these systems (e.g., Bethke, 1984; Rye, 1993; Hedenquist and Lowenstern, 1994; Arribas, 1995; Cooke and Simmons, 2000). This magmatic component originates from the exsolution of a fluid from the melt during retrograde (or resurgent) boiling (Burnham, 1979; Burnham and Ohmoto, 1980). When the exsolved fluid reaches its solvus, the inception of aqueous immiscibility leads to the formation of low-density vapor and dense hypersaline liquid. This process is critical for the partitioning of ore metals from the melt to the hydrothermal system (Candela and Piccoli, 1995; Williams et al., 1995; Heinrich et al., 1999). The vapor phase will rise to shallow depths and condense, and the ensuing low-pH

fluids will react with wallrocks. Spent fluids will either discharge at surface as volcanic fumaroles, or be absorbed into groundwater. The dense, hypersaline liquid remains at depth and is recorded as saline fluid inclusions in intrusive rocks and porphyry ore deposits (Hedenquist and Lowenstern, 1994 and references therein), or becomes part of later epithermal mineralization (Rye, 1993). As magmatic fluids rise through the crust, they typically undergo phase separation at the transition from brittle to plastic behavior in the lithosphere (Fournier, 1999). This transition typically occurs at about 370 to 400 °C in continental hydrothermal systems (Fournier, 1999). Abrupt changes in the pressure regime across this transition zone are thought to be responsible for porphyry Cu mineralization in several systems (e.g., Fournier, 1967; Gustafson and Hunt, 1975; Cunningham, 1978), and for porphyry Au mineralization (Muntean and Einaudi, 2000). Similarly, several authors have related epithermal mineralization to the escape of magmatic fluids into hydrostatically pressured hydrothermal systems (e.g., Fournier, 1987; Rye, 1993; Deen et al., 1994; Hedenquist et al., 1994). In the following discussion, we examine the role of magmatic fluids in the Tambo high-sulfidation system, and the nature of magmatic-steam processes.

9.1. Role of magmatic fluids in alunite deposition

It is generally accepted that hypogene acid-sulfate alteration, characterized by the assemblage alunite+quartz+kaolinite±pyrite, results from acidic fluids that are generated by the condensation of magmatic volatiles enriched in SO₂, HCl, and HF. The disproportionation of SO₂, beginning at about 400 °C, and the dissociation of HCl and HF at temperatures below about 300–350 °C (Holland, 1965; Hemley et al., 1969) create solutions that are sufficiently acidic to leach most components from the host rock, leaving a vuggy quartz residue. The progressive neutralization of these fluids by reaction with wallrock forms successive alteration envelopes of acid-sulfate, advanced argillic, argillic, and propylitic assemblages outward from the main fluid conduit (e.g., Steven and Ratté, 1960). Stable-isotope evidence indicates that the degree of meteoric water interaction during hypogene acid-sulfate alteration is variable. At the Julcani deposit in Peru, for example, the condensed

liquids responsible for magmatic hydrothermal alunite were almost entirely of magmatic origin (Deen et al., 1994). A large component of magmatic fluid was also present at Rodalquilar (Arribas et al., 1995). By contrast, the fluids responsible for magmatic hydrothermal alunite in many other high-sulfidation deposits had a much more extensive component of meteoric water (e.g., Summitville, Rye et al., 1990; Nansatsu, Hedenquist et al., 1994; Pueblo Viejo, Venneman et al., 1993).

Examples of magmatic contributions to the near-surface steam-heated environment are fewer. Steam-heated alunite forms from aqueous sulfate created by the oxidation of H₂S at or above the water table (Schoen et al., 1974). The fluids responsible for this alunite typically have a meteoric isotopic signature (e.g., Rye et al., 1992; Ebert and Rye, 1997); however, deuterium enrichment (10–20‰) is common in steam-heated fluids from active geothermal systems. This enrichment can be attributed to the effects of either subsurface boiling or evaporation (Truesdell et al., 1977; Henley and Stewart, 1983). However, magmatic fluids with low δD values have been documented in shallow steam-heated environments (Rye et al., 1992).

Shallow late-stage alteration at Tambo also involved a large magmatic fluid component. Stable isotopic data indicate that the parental fluids responsible for this late-stage alteration were mixtures ranging from nearly pure magmatic fluids with low δD (either magmatic steam or condensed magmatic vapor) to dominantly meteoric water. The variability in trace-element contents for alunite of this stage is attributed to different magmatic and meteoric fluid mixtures in the shallow epithermal environment.

Overall, there is a strong magmatic fluid component to parental fluids for all alunite types at the Tambo deposit, even in near-surface and late-stage occurrences. Even the groundwater at Tambo seems to have had a substantial component of condensed magmatic fluid. These fluids effectively displaced or mixed with meteoric groundwaters. We attribute the dominance of magmatic fluids at Tambo to climatic and physiographic characteristics specific to the El Indio–Pascua belt, as detailed in Bissig (2001) and Deyell et al. (2004). The region was dominated by extensive sub-planar surfaces that were incised during regional erosion starting in the late Miocene (Bissig, 2001). The near-surface fluid dynamics in this setting

(Bissig, 2001), combined with arid conditions, presumably limited the availability of meteoric water in the upper part of the Tambo system.

9.2. Role of magmatic fluids in mineralization

Although the consensus is that the fluids responsible for high-sulfidation deposits are of magmatic origin (e.g., Rye et al., 1992; Hedenquist and Lowenstern, 1994; Arribas, 1995; Cooke and Simmons, 2000), there is considerable debate about the nature of these fluids, and particularly about the relative contributions of vapor versus hypersaline liquid to the formation of Au, Ag, and Cu mineralization. Several authors have argued that mineralization is a two-stage process involving both vapor and later saline magmatic fluids (e.g., Stoffregen, 1987; Rye, 1993; Arribas, 1995). On the basis of the results of fluid-inclusion and isotopic studies of several high-sulfidation deposits, three possible ore-fluid types were summarized by Cooke and Simmons (2000):

- (a) Low-salinity, supercritical magmatic fluids which cool to a single-phase liquid before boiling at shallow levels and which transport Au and possibly Cu as H₂S complexes (Loucks and Mavrogenes, 1999).
- (b) Metal-bearing, acidic brines, derived from the second boiling of magmatic fluids, transport Au and Cu presumably largely as chloride complexes (Henley, 1973).
- (c) Reduced magmatic vapors (the counterpart of the brines from the second boiling of magmatic fluids) that transport Au and Cu as volatile sulfide, bisulfide, or chloride complexes (Henley, 1973; Seward, 1973; Gibert et al., 1998).

Recently, Muntean and Einaudi (2001) proposed that high-sulfidation gold ore at La Pepa (Maricunga belt) was formed simultaneously with quartz-alunite alteration by the condensation of a moderately saline, supercritical fluid that was later flashed, from below the brittle–ductile transition, into shallow meteoric waters.

Recent studies of metal fractionation between magmatic brines and vapors, and evidence from fluid and melt inclusions, indicate that magmatic vapors have the capacity to transport metals such as Cu and Au

for high-sulfidation mineralization, particularly at high pressures (Lowenstern et al., 1991; Heinrich et al., 1992; Shinohara, 1994; Audétat et al., 1998; Heinrich et al., 1999; Ulrich et al., 1999; Archibald et al., 2001). At Tambo, Stage 2 alunite and ore resulted from H₂S-dominant, low-salinity magmatic vapor (this study; Jannas et al., 1999). There is no evidence for a hypersaline fluid component; nearly all fluid inclusions measured by Jannas (1995) contain <4 wt.% NaCl equivalent. We infer that Au was transported in a magmatic vapor phase, either as volatile species (e.g., AuS_(g)) or as bisulfide complexes. The high Te content of the mineralizing assemblage is also consistent with vapor-phase transport (Cooke and McPhail, 2000). The Stage 2 high-sulfidation mineralization most likely precipitated from condensed magmatic vapor that rose from the brittle–ductile transition and was responsible for the earlier acid-sulfate alteration.

The nature of Stage 3 ore fluids is less clear, but again there is no fluid-inclusion or gas-chemistry evidence for a hypersaline brine component. Stage 3 fluids were more vapor-rich magmatic fluids and more oxidized (SO₂-rich) than those responsible for Stage 2 Au deposition. There are two viable scenarios for these fluids: (1) they may have been derived as vapor directly from a magma, or (2) they may have been derived as flashed liquids from below the brittle–ductile transition (much as suggested by Muntean and Einaudi, 2001). The latter scenario is suggested by the gas-chemistry data, in particular the lack of He in fluid inclusions within the alunite of magmatic-steam origin (Landis and Rye, *this volume*) and the lack of sawtooth banding in the trace-element pattern of this type of alunite. Presumably, some Au was transported in these fluids as a chloride complex (Archibald et al., 2001).

9.3. Origin of magmatic-steam alunite

The magmatic-steam environment was initially defined by Rye et al. (1992) on the basis of data for veins of coarsely crystalline alunite at Marysvale, Utah (Cunningham et al., 1984). This alunite is characterized by disequilibrium $\delta^{34}\text{S}$ but near-equilibrium $\delta^{18}\text{O}$ and magmatic δD values. Documented occurrences of this type of alunite are rare, and magmatic-steam processes are poorly understood. Magmatic steam was important at the Tambo deposit, however, because of its abundance and association with Stage 3 ore. Stable-

isotope and fluid-inclusion data from this study and that of Landis and Rye (this volume) help to characterize the magmatic-steam environment. Important observations are summarized below.

- (a) Like the Marysvale magmatic-steam alunite, the inclusion fluids for the Tambo alunite have much lower $\text{H}_2\text{S}/\text{SO}_2$ than inclusion fluids in magmatic-hydrothermal alunite.
- (b) Alunite of magmatic-steam origin at Tambo is also characterized by having nearly constant $\delta^{34}\text{S}$ values that are close to the value of the bulk S in the magma (Rye et al., 1992). Given that sulfur-isotope equilibrium between aqueous sulfate and sulfide is obtained quickly at high temperature (Ohmoto and Lasaga, 1982), a rapid rate of vapor ascent and mineral deposition is required once the fluids leave their source (Rye et al., 1992; Rye, 1993).
- (c) Inclusion fluids of both the Marysvale and Tambo magmatic-steam alunite are vapor-rich.
- (d) Tambo magmatic-steam alunite is significantly enriched in LREE and strongly fractionated in HREE compared with the host rock. These data may reflect REE partitioning into the vapor phase (e.g., Flynn and Burnham, 1978; Reed, 1995).
- (e) Magmatic-steam alunite formed late in the development of the Tambo hydrothermal system. A similar transition from early magmatic-hydrothermal to late magmatic-steam processes has also been documented at the Red Mountain deposit, Colorado (Bove et al., 1990; Rye et al., 1992).
- (f) Two important differences between the Marysvale and Tambo magmatic-steam alunite are that the fluid inclusions at Tambo are depleted in He and they do not contain the sawtooth trace-element patterns observed for Marysvale alunite. This pattern for Marysvale alunite is thought to be related directly to magmatic degassing (Cunningham et al., 1996; Rye, this volume; Landis and Rye, this volume). The lack of He in the Tambo magmatic-steam alunite indicates that the magmatic vapor was already degassed and thus could not have come directly from a melt.

The parental fluids for the magmatic-steam alunite at Tambo were most likely derived by flashing, due to sudden decompression, of condensed magmatic fluids

that accumulated below the brittle–ductile transition near a shallow magma. This scenario is similar to that proposed by Muntean and Einaudi (2001) for high-level Au mineralization at La Pepa, Chile. After decompression of the system, the ensuing vapor fluids were SO_2 -dominant because of the effect of high temperature and low pressure on the gas equilibria of the sulfur species (Gerlach, 1993). Prior loss of H_2S to the vapor phase may have also increased the oxidation state of the residual liquids.

10. Summary

Detailed argon-age, stable-isotope, fluid-inclusion gas, and geochemical analyses of alunite from the Tambo deposit, coupled with the geological framework determined by Jannas et al. (1999), have allowed a study of shallow-level magmatic-hydrothermal processes. Hydrothermal activity at Tambo was characterized by multiple stages of alteration and two mineralizing events related to episodic magmatic activity over a period of at least 4 my. We have focused on the characteristics of alunite within this deposit to gain insight into the nature of parental fluids in magmatic-hydrothermal, magmatic-steam, steam-heated, and apparent supergene environments. Results from this study have shown:

- (1) There is a strong magmatic signature to all stages and environments of acid alteration in the Tambo area, even in the near-surface steam-heated environment and at shallow levels during the final stages of the hydrothermal system, in what is easily mistaken for supergene alteration. The meteoric water component in fluids of all stages of alteration is ^{18}O -enriched. The isotopic data for fluids reflect unusually arid conditions during mineralization.
- (2) Magmatic vapor-phase transport of Au is inferred for both Stage 2 and Stage 3 mineralization.
- (3) The fluids responsible for both alteration and the associated Stage 2 and 3 mineralization evolved from high-temperature magmatic vapor-bearing fluids that had different origins for each stage. The vapor responsible for the deposition of magmatic-hydrothermal alunite and Stage 2 mineralization was derived as a long-standing, H_2S -rich plume that rose above the brittle–

ductile transition. This vapor was derived from the “second boiling” of the magmatic fluids. The vapor responsible for magmatic-steam alunite and Stage 3 mineralization was SO₂-rich and probably was derived from the flashing of partly degassed and oxidized saline fluids beneath the brittle–ductile transition. The transition from magmatic-hydrothermal to magmatic-steam-dominated processes probably reflects sudden decreases in pressure in the hydrothermal system.

Acknowledgements

This paper stems from a PhD project undertaken by the lead author at the University of British Columbia. Funding was provided by Barrick Gold and Barrick Chile, with additional support from NSERC. This study would not have been possible without the expertise and knowledge of Barrick geologists, particularly R. Guerra and the exploration staff at Km. 34, and the Tambo mine staff. Discussions and assistance provided by K. Heather, D. Heberlein, N. Rojas, and R. Jannas are greatly appreciated. PIMA instrumentation and expertise was provided by A. Thompson and Petrascience Consulting. C. Kester and M. Raudsepp are thanked for their technical assistance. This manuscript benefited greatly from early reviews and comments by S. Ebert, J. Thompson, J. Hedenquist, P. Bethke, A. Hofstra, L. Groat, and J. Richards. We also thank the *Chemical Geology* reviewers, K. Heather and G. Arehart, and editors for their constructive comments. [PD]

References

- Aoki, M., 1991. Mineralogical features and genesis of alunite solid solution in high temperature magmatic-hydrothermal systems. *J. Geol. Surv. Jpn.* 277, 31–32.
- Aoki, M., Comsti, E.C., Lazo, F.B., Matsuhisa, Y., 1993. Advanced argillic alteration and geochemistry of alunite in an evolving hydrothermal system at Baguio, northern Luzon, Philippines. *Resour. Geol.* 43, 155–164.
- Archibald, S.M., Migdisov, A.A., Williams-Jones, A.E., 2001. The stability of Au-chloride complexes in water vapor at elevated temperature and pressures. *Geochim. Cosmochim. Acta* 65, 4413–4423.
- Arribas Jr., A., 1995. Characteristics of high sulfidation epithermal deposits, and their relation to magmatic fluids. In: Thompson, J.F.H. *Magmas, Fluids, and Ore Deposits*. Mineral. Assoc. Can. Short Course vol. 23. , pp. 419–454.
- Arribas Jr., A., Cunningham, C.G., Rytuba, J.J., Rye, R.O., Kelly, W.C., McKee, E.H., Podwysocny, M.H., Tosdal, R.M., 1995. Geology, geochronology, fluid inclusions, and stable isotope geochemistry of the Rodalquilar Au alunite deposit, Spain. *Econ. Geol.* 90, 795–822.
- Audétat, A., Günther, D., Heinrich, C.A., 1998. Formation of magmatic-hydrothermal ore deposit: Insights with LA-ICP-MS analysis of fluid inclusions. *Science* 279, 2091–2094.
- Barazangi, M., Isacks, B.L., 1976. Spatial distribution of earthquakes and subduction of the Nazca Plate beneath South America. *Geology* 4, 686–692.
- Beane, R.E., 1991. Deposit correlations in the El Indio Belt, Chile: mineral paragenesis, fluid inclusions, and sulfur isotopes. Barrick Gold and Barick Chile, Internal company report, La Serena, Chile.
- Bennet, R.E., 1995. A combined hydrothermal alteration study and exploration target evaluation of the Tambo mining district, El Indio mineral belt, Region IV, Chile. Compañía Minera San José, Internal company report, Chile.
- Bethke, P.M., 1984. Controls on base- and precious-metal mineralization in deeper epithermal environments. *U.S. Geol. Surv. Open-File Rep.*, 84–890.
- Bissig, T., 2001. Metallogenesis of the Miocene El Indio-Pascua gold-silver-copper belt, Chile/Argentina: Geodynamic, geomorphological and petrochemical controls on epithermal mineralization. Ph.D. thesis, Queen’s University, Kingston, Canada.
- Bove, D.J., Rye, R.O., Hon, K., 1990. Evolution of the Red Mountain alunite deposits, Lake City, Colorado. *U.S. Geol. Surv. Open-File*, 90–102.
- Burnham, C.W., 1979. Magmas and hydrothermal fluids. In: Barnes, H.L. (Ed.), *Geochemistry of Hydrothermal Ore Deposits*. Wiley, New York, pp. 71–136.
- Burnham, W.C., Ohmoto, H., 1980. Late-stage processes of felsic magmatism. *Min. Geol. Spec. Issue* 8, 1–11.
- Candela, P.A., Piccoli, P.M., 1995. Model ore-metal partitioning from melts into vapor and vapor–brine mixtures. In: Thompson, J.F.H. (Ed.), *Magmas, Fluids, and Ore Deposits*, Mineral. Assoc. Can. Short Course vol. 23. , pp. 101–127.
- Clayton, R.N., Mayeda, T.K., 1963. The use of bromine pentafluoride in the extraction of oxygen from oxides and silicates for isotopic analysis. *Geochim. Cosmochim. Acta* 27, 43–52.
- Cooke, D.R., McPhail, D.C., 2000. Epithermal Au–Ag–Te mineralization, Acupan, Baguio District, Philippines: Numerical simulations of mineral deposition. *Econ. Geol.* 96, 109–131.
- Cooke, D.R., Simmons, S.F., 2000. Characteristics and genesis of epithermal gold deposits. In: Hagemann, S.G., Brown, P.E. (Eds.), *Gold 2000*, Rev. *Econ. Geol.* vol. 13, pp. 221–244.
- Craig, H., 1961. Isotopic variations in meteoric waters. *Science* 133, 1702–1703.
- Craig, H., Lupton, J.E., 1981. ³He and mantle volatiles in the ocean and the ocean crust. In: Emiliani, C. (Ed.), *The Sea; Ideas and Observations on Progress in the Study of the Seas*. Wiley, New York, pp. 391–428.

- Cunningham, C.G., 1978. Pressure gradients and boiling as mechanisms for localizing ore in porphyry systems. *U.S. Geol. Surv. J. Res.* 6, 745–754.
- Cunningham, C.G., Rye, R.O., Steven, T.A., Mehnert, H.H., 1984. Origins and exploration significance of replacement and vein-type alunite deposits in the Marysvale volcanic field, west central Utah. *Econ. Geol.* 79, 50–71.
- Cunningham, C.G., Rye, R.O., Bethke, P.M., Logan, M.A.V., 1996. Formation of coarse-grained vein alunite by degassing of an epizonal stock. Chapman Conference on Crater Lakes, Terrestrial Degassing and Hyper-Acid Fluids in the Environment, Extended Abstracts. Sept. 4–9, 1996, Crater Lake, Oregon. *Am. Geophys. Union, Washington, DC*, pp. 27.
- Deen, J.A., Rye, R.O., Munoz, J.L., Drexler, J.W., 1994. The magmatic-hydrothermal system at Julcani, Peru: evidence from fluid inclusions and hydrogen and oxygen isotopes. *Econ. Geol.* 89, 1924–1938.
- Deyell, C.L., 2001. Alunite and high sulfidation Au–Ag–Cu mineralization in the El Indio–Pascua Belt, Chile–Argentina, PhD thesis, Univ. British Columbia, Vancouver, Canada.
- Deyell, C.L., Bissig, T., Rye, R.O., 2004. Isotopic evidence for magmatic-dominated epithermal processes in the El Indio–Pascua Au–Cu–Ag belt and relationship to geomorphologic settings. *Econ. Geol. Spec. Publ.* 11, 55–73.
- Ebert, S.W., Rye, R.O., 1997. Secondary precious metal enrichment by steam-heated fluids in the Crowfoot–Lewis hot spring gold–silver deposit and relation to paleoclimate. *Econ. Geol.* 92, 578–600.
- Flynn, F.R., Burnham, C.W., 1978. An experimental determination of rare earth partition coefficients between a chloride-containing vapor phase and silicate melts. *Geochim. Cosmochim. Acta* 42, 685–701.
- Fournier, R.O., 1967. The porphyry copper deposit exposed in the Liberty open-pit mine near Ely, Nevada: Part I. Syngenetic formation. *Econ. Geol.* 62, 57–81.
- Fournier, R.O., 1987. Conceptual models of brine evolution in magmatic-hydrothermal systems. *U.S. Geol. Surv. Prof. Pap.* 1350 (2), 1487–1506.
- Fournier, R.O., 1999. Hydrothermal processes related to movement of fluid from plastic into brittle rock in the magmatic-hydrothermal environment. *Econ. Geol.* 94, 1193–1212.
- Fulignati, P., Gioncada, A., Sbrana, A., 1999. Rare-earth element behaviour in the alteration facies of the active magmatic-hydrothermal systems of Vulcano (Aeolian Islands, Italy). *J. Volcanol. Geotherm. Res.* 88, 325–342.
- Gerlach, T.M., 1993. Oxygen buffering of Kilauea volcanic gases and the oxygen fugacity of Kilauea basalt. *Geochim. Cosmochim. Acta* 57, 795–814.
- Gerling, E.K., Mamyrin, B.A., Tolstikhin, I.N., 1971. Helium isotope composition in some rocks. *Geochem. Int.* 8, 755–761.
- Gieseman, A., Jager, H.J., Norman, A.L., Krouse, H.R., Brand, W.A., 1994. On-line sulfur-isotope determination using an elemental analyzer coupled to a mass spectrometer. *Anal. Chem.* 66, 2816–2819.
- Gibert, F., Pascal, M.L., Pichavant, M., 1998. Gold solubility and speciation in hydrothermal solutions: Experimental study of the stability of hydrosulfide complex of gold (AuHS^o) at 350 to 450 °C and 500 bars. *Geochim. Cosmochim. Acta* 62, 2931–2947.
- Giggenbach, W.F., 1992. Magma degassing and mineral deposition in hydrothermal systems along convergent plate boundaries. *Econ. Geol.* 87, 1927–1944.
- Godfrey, J.D., 1962. The deuterium content of natural waters and other substances. *Geochim. Cosmochim. Acta* 27, 43–52.
- Gordon, S., McBride, B.J., 1994. Computer program for calculation of complex chemical equilibrium compositions and applications: I. Analysis. NASA Ref. Publ. 1311, 55 pp.
- Graney, J.R., Kesler, S.E., 1995a. Gas composition of inclusion fluid in ore deposits: is there a relation to magmas? In: Thompson, J.F.H. (Ed.), *Magmas, Fluids, and Ore Deposits*, Mineral. Assoc. Can. Short Course, vol. 23, pp. 221–246.
- Graney, J.R., Kesler, S.E., 1995b. Factors affecting gas analysis of inclusion fluid by quadrupole mass spectrometry. *Geochim. Cosmochim. Acta* 59, 3977–3986.
- Guha, J., Lu, L.Z., Gagnon, M., 1990. Gas composition of fluid inclusions using solid probe mass spectrometry and its application to study of mineralizing processes. *Geochim. Cosmochim. Acta* 54, 553–558.
- Gustafson, L.B., Hunt, J.P., 1975. The porphyry copper deposit at El Salvador, Chile. *Econ. Geol.* 70, 857–912.
- Heather, K.B., Diaz, D., 2000. El Indio–Tambo District regional geological mapping program 1997–1999, Barick Chile, Internal company report, 2 Vol., La Serena, Chile.
- Hedenquist, J.W., Lowenstern, J.B., 1994. The role of magmas in the formation of hydrothermal ore deposits. *Nature* 370, 519–527.
- Hedenquist, J.W., Matsuhisa, Y., Izawa, E., White, N.C., Giggenbach, W.F., Aoki, M., 1994. Geology, geochemistry, and origin of high sulfidation Cu–Au mineralization in the Nansatsu District, Japan. *Econ. Geol.* 89, 1–30.
- Heinrich, C.A., Ryan, C.G., Mernagh, T.P., Eadington, P.J., 1992. Segregation of ore metals between magmatic brine and vapor: a fluid inclusion study using PIXE microanalysis. *Econ. Geol.* 87, 1566–1583.
- Heinrich, C.A., Gunther, D., Audétat, A., Ulrich, T., Frischknecht, R., 1999. Metal fractionation between magmatic brine and vapor, determined by microanalysis of fluid inclusions. *Geology* 27, 755–758.
- Hemley, J.J., Hostetler, P.B., Gude, A.J., Mountjoy, W.T., 1969. Some stability relations of alunite. *Econ. Geol.* 64, 599–612.
- Henley, R.W., 1973. Solubility of gold in hydrothermal chloride solutions. *Chem. Geol.* 11, 73–87.
- Henley, R.W., Stewart, M.K., 1983. Chemical and isotopic changes in the hydrology of the Tauhara geothermal field due to exploitation at Wairakei. *J. Volcanol. Geotherm. Res.* 15, 285–314.
- Holland, H.D., 1965. Some applications of thermochemical data to problems of ore deposits: II. Mineral assemblages and the compositions of ore-forming fluids. *Econ. Geol.* 60, 1101–1166.
- Hopf, S., 1993. Behavior of rare earth elements in geothermal systems of New Zealand. *J. Geochem. Explor.* 47, 333–357.
- Jambor, J.L., 1999. Nomenclature of the alunite supergroup. *Can. Mineral.* 37, 1323–1341.

- Jannas, R.R., 1995. Reduced and oxidized high sulfidation deposits of the El Indio deposit, Chile. PhD thesis, Harvard Univ., Cambridge, MA.
- Jannas, R.R., Bowers, T.S., Petersen, U., Beane, R.E., 1999. High-sulfidation deposit types in the El Indio District, Chile. In: Skinner, B.J. (Ed.), *Geology and Ore Deposits of the Central Andes*, Soc. Econ. Geol. Spec. Pub., vol. 7, pp. 27–59.
- Kennedy, B.M., Hiyagon, H., Reynolds, J.H., 1990. Crustal neon: a striking uniformity. *Earth Planet. Sci. Lett.* 98, 277–286.
- Kornexl, B.E., Gehre, M., Höfling, R., Werner, R.A., 1999. On-line $\delta^{18}\text{O}$ measurement of organic and inorganic substances. *Rapid Commun. Mass Spectrom.* 13, 1685–1693.
- Landis, G.P., Hofstra, A.H., 1991. Fluid inclusion gas chemistry as a potential mineral exploration tool: Case studies from Creede, CO, Jerrit Canyon, NV, Coeur d'Alene district, ID and MT, southern Alaska epithermal veins, and mid-continent MVT's. *J. Geochem. Explor.* 42, 25–60.
- Landis, G.P., Rye, R.O., 2005. Characterization of gas chemistry and noble-gas isotopic ratios of inclusion fluids in magmatic-hydrothermal environments. *Chem. Geol.* 215, 155–184. (this volume).
- Lewis, A.J., Palmer, M.R., Sturchio, N.C., Kemp, A.J., 1997. The rare earth element geochemistry of acid-sulphate and acid-sulphate-chloride geothermal systems from Yellowstone National Park, Wyoming, USA. *Geochim. Cosmochim. Acta* 61, 695–706.
- Lewis, A.J., Komminou, A., Yardley, B.W.D., Palmer, M.R., 1998. Rare earth element speciation in geothermal fluids from Yellowstone National Park, Wyoming, USA. *Geochim. Cosmochim. Acta* 62, 657–663.
- Li, G., Peacor, D.R., Essene, E.J., Brosnahan, D.R., Beane, R.E., 1992. Walthierite and huangite, two new minerals of the alunite group from the Coquimbo region, Chile. *Am. Mineral.* 77, 1275–1284.
- Lindaas, S.E., Chomiak, B.A., Campbell, A.R., Norman, D.I., 1998. Enargite-hosted fluid inclusion gases from the Lepanto, Philippines, high-sulfidation Cu–Au deposit. *Geol. Soc. Am. Abstr. Programs* 30 (7), 20.
- Lottermoser, B.J., 1992. Rare earth elements and hydrothermal ore formation processes. *Ore Geol. Rev.* 7, 25–41.
- Loucks, R.R., Mavrogenes, J.A., 1999. Gold solubility in supercritical hydrothermal brines measured in synthetic fluid inclusions. *Science* 284, 2159–2163.
- Lowenstern, J.B., Mahood, G.A., Rivers, M.L., Sutton, S.R., 1991. Evidence for extreme partitioning of copper into a magmatic vapor phase. *Science* 257, 1405–1409.
- Lupton, J.E., 1983. Terrestrial inert gases: isotope tracer studies and clues to primordial components in the mantle. *Ann. Rev. Earth Sci.* 11, 371–414.
- Maksaev, V., Moscoso, R., Mpodozis, C., Nasi, C., 1984. Las unidades volcánicas y plutónicas del cenozoico superior en la Alta Cordillera del Norte Chico (29°–31° S): Geología, alteración hidrotermal y mineralización. *Rev. Geol. Chile* 21, 11–51.
- Mamyrin, B.A., Anuffriev, G.S., Kamenski, I.L., Tolstikhin, I.N., 1969. Determination of the isotopic composition of atmospheric helium. *Geochem. Int.* 7, 498–505.
- Martin, M., Clavero, J., Mpodozis, C., 1995. Estudio geológico regional del Franja El Indio, Cordillera de Coquimbo. Servicio Nacional de Geología y Minería, Santiago, Chile. Registered report IR-95-06, 232 pp.
- Martin, M.W., Clavero, J.R., Mpodozis, C.M., 1997. Eocene to Late Miocene magmatic development of the El Indio belt, ~30 °S, north-central Chile. *Congreso Geol. Chileno, VIII*, Concepción Chile, Actas vol. 1. , pp. 149–153.
- Martin, M.W., Clavero, J.R., Mpodozis, C., 1999. Late Paleozoic to Early Jurassic tectonic development of the high Andean Principal Cordillera, El Indio Region, Chile (29–30°S). *J. South Am. Earth Sci.* 12, 33–49.
- Marty, B., 1989. Neon and xenon isotopes in MORB: implications for the earth-atmosphere evolution. *Earth Planet. Sci. Lett.* 94, 45–56.
- McBride, B.J., Gordon, S., 1996. Computer program for calculation of complex chemical equilibrium compositions and applications: II. Users manual and program description. NASA Ref. Publ., 1311.
- Michard, A., 1989. Rare earth element systematics in hydrothermal fluids. *Geochim. Cosmochim. Acta* 53, 745–750.
- Morrison, P., Pine, J., 1955. Radiogenic origin of the helium isotopes in rock. *N.Y. Acad. Sci., Ann.* 62, 71–92.
- Muntean, J.L., Einaudi, M.T., 2000. Porphyry gold deposits of the Refugio district, Maricunga belt, northern Chile. *Econ. Geol.* 95, 1445–1472.
- Muntean, J.L., Einaudi, M.T., 2001. Porphyry-epithermal transition: Maricunga belt, northern Chile. *Econ. Geol.* 96, 743–772.
- Ohmoto, H., Lasaga, A.C., 1982. Kinetics of reactions between aqueous sulfates and sulfides in hydrothermal systems. *Geochim. Cosmochim. Acta* 46, 1727–1746.
- Ozima, M., Podosek, F.A., 2002. *Noble Gas Geochemistry*. Cambridge Univ. Press, Cambridge, UK.
- Petersen, E.U., Thompson, A.J.B., 1992. Analysis of alunite and other thermally sensitive minerals. *Microprobe Analyst* (Internal report, Dep. Geol. Geophys., Univ. Utah), April, pp. 2–6.
- Pickthorn, W.J., O'Neil, J.R., 1985. ^{18}O relations in alunite mineral: potential single mineral thermometer. *Geol. Soc. Am. Abstr. Programs* 17 (7), 686.
- Poreda, R., Radicati di Brozolo, F., 1984. Neon in mid-Atlantic ridge basalts. *Earth Planet. Sci. Lett.* 69, 277–289.
- Pouchou, J.L., Pichoir, F., 1985. PAP (ρZ) procedure for improved quantitative microanalysis. *Microbeam Anal.* 1985, 104–106.
- Ramos, V.A., Mahlburg, K.S., Page, R.N., Munizaga, F., 1989. La ignimbrita Vacas Heladas y el cese del volcanismo en el Valle del Cura, Provincia de San Juan. *Rev. Asoc. Geol. Argent.* 44, 336–352.
- Reed, M.J., 1995. Distribution of rare earth elements between aqueous fluids and granitic melt, PhD thesis, Univ. Maryland at College Park.
- Rye, R.O., 1993. The evolution of magmatic fluids in the epithermal environment: the stable isotope perspective. *Econ. Geol.* 88, 733–753.
- Rye, R.O., 2005. A review of the stable-isotope geochemistry of sulfate minerals in selected igneous environments and related hydrothermal systems. *Chem. Geol.* 215, 5–36. (this volume).
- Rye, R.O., Stoffregen, R.E., Bethke, P.M., 1990. Stable isotope systematics and magmatic hydrothermal processes in the

- Summitville, CO, gold deposit. U.S. Geol. Surv. Open-File, 90–626.
- Rye, R.O., Bethke, P.M., Wasserman, M.D., 1992. The stable isotope geochemistry of acid sulfate alteration. *Econ. Geol.* 87, 225–262.
- Sandeman, H.A., Archibald, D.A., Grant, J.W., Villeneuve, M.E., Ford, F.D., 1999. Characterisation of the chemical composition and ^{40}Ar – ^{39}Ar systematics of intralaboratory standard MAC-83 biotite. Radiogenic age and isotopic studies: report 12. *Geol. Surv. Can. Current Res.* 1999-F, 13–26.
- Savin, S.M., Epstein, S., 1970. The oxygen and hydrogen isotope geochemistry of clay minerals. *Geochim. Cosmochim. Acta* 34, 24–42.
- Schoen, R., White, D., Hemley, J.J., 1974. Argillization by descending acid at steamboat springs, Nevada. *Clays Clay Miner.* 22, 1–22.
- Schwab, R.G., Herold, H., Gotz, C., de Oliveira, N.P., 1990. Compounds of the crandallite type: synthesis and properties of pure rare earth element phosphates. *Neues Jahrb. Mineral.*, 241–254.
- Seward, T.M., 1973. Thio complexes of gold and the transport of gold in hydrothermal ore solutions. *Geochim. Cosmochim. Acta* 37, 379–399.
- Shepherd, T.J., Rankin, A.H., 1998. Fluid inclusion techniques of analysis. In: Richards, J.P., Larson, P.B. (Eds.), *Techniques in Hydrothermal Ore Deposits Geology*, Rev. *Econ. Geol.*, vol. 10, pp. 125–150.
- Shinohara, H., 1994. Exsolution of immiscible vapor and liquid phases from a crystallizing silicate melt: Implications for chlorine and metal transport. *Geochim. Cosmochim. Acta* 58, 5215–5221.
- Siddleley, G., Areneda, R., 1985. Gold–silver occurrences of the El Indio belt, Chile. In: Ericksen, G.E., Pinochet, M.T., Reinemund, J.A. (Eds.), *Geology of the Andes and Its Relation to Hydrocarbon and Mineral Resources*. Circum-Pacific Council for Energy and Mineral Resources, Earth Science Series, pp. 273–284.
- Sillitoe, R.H., 1999. Styles of high sulfidation gold, silver and copper mineralization in the porphyry and epithermal environments. PACRIM '99 congress. Bali Indonesia, 10–13 October, Proc.-Australas. Inst. Min., pp. 29–44.
- Steiger, R.H., Jäger, E., 1977. Subcommittee on geochronology: convention on the use of decay constants in geo- and cosmochronology. *Earth Planet. Sci. Lett.* 36, 359–362.
- Steven, T.A., Ratté, J.C., 1960. Geology of ore deposits of the Summitville district, San Juan Mountains, Colorado. *U.S. Geol. Surv. Prof.*, 343.
- Stoffregen, R.E., 1987. Genesis of acid-sulfate alteration and Au–Cu–Ag mineralization at Summitville, Colorado. *Econ. Geol.* 82, 1575–1591.
- Stoffregen, R.E., Alpers, C.N., 1987. Woodhouseite and svanbergite in hydrothermal ore deposits: products of apatite destruction during advanced argillic alteration. *Can. Mineral.* 25, 201–211.
- Stoffregen, R.E., Cygan, G., 1990. An experimental study of Na–K exchange between alunite and aqueous sulfate solutions. *Am. Mineral.* 75, 209–220.
- Stoffregen, R.E., Rye, R.O., Wasserman, M.D., 1994. Experimental studies of alunite: I. ^{18}O – ^{16}O and D–H fractionation factors between alunite and water at 250–450 °C. *Geochim. Cosmochim. Acta* 58, 903–916.
- Taylor, B.E., 1986. Magmatic volatiles: isotopic variation of C, H, S. In: Valley, J.W., Taylor Jr., H.P., O'Neil, J.R. (Eds.), *Stable Isotopes in High Temperature Geological Processes*, Rev. Mineral., vol. 16, pp. 185–225.
- Taylor, B.E., 1988. Degassing of rhyolitic magmas: hydrogen isotope evidence and implications for magmatic-hydrothermal ore deposits. *Can. Inst. Min. Metall. Spec. vol.* 39, 33–49.
- Taylor, B.E., 1991. Degassing of Obsidian Dome rhyolite, Inyo volcanic chain, California. In: Taylor Jr., H.P., O'Neil, J.R., Kaplan, I.R. (Eds.), *Stable Isotope Geochemistry: A Tribute to Samuel Epstein*, *Geochem. Soc. Spec. Pub.*, vol. 3, 339–353.
- Thompson, A.J.B., 1992. Alunite compositions and textures: relationships to precious metal mineralization. Mineral Deposit Research Unit Short Course #8: New Developments in Litho-geochemistry. University of British Columbia, Vancouver, Canada.
- Thompson, A.J.B., Hauff, P.L., Robitaille, A.J., 1999. Alteration mapping in exploration: Application of short-wave infrared (SWIR) spectroscopy. *Soc. Econ. Geol. Newsl.* 39, 13. (Oct.).
- Truesdell, A.H., Nathenson, M., Rye, R.O., 1977. The effects of boiling and dilution on the isotopic compositions of Yellowstone thermal waters. *J. Geophys. Res.* 82, 3694–3704.
- Ulrich, T., Gunther, D., Heinrich, C.A., 1999. Gold concentrations of magmatic brines and the metal budget of porphyry copper deposits. *Nature* 399, 676–679.
- Venneman, T.W., Muntean, J.L., Kesler, S.E., O'Neil, J.R., Valley, J.W., Russell, N., 1993. Stable isotope evidence for magmatic fluids in the Pueblo Viejo epithermal acid sulfate Au–Ag deposit, Dominican Republic. *Econ. Geol.* 88, 55–71.
- Wasserman, M.D., Rye, R.O., Bethke, P.M., Arribas Jr., A., 1992. Methods for separation and total stable isotope analysis of alunite. *U.S. Geol. Survey. Open-File Rep.*, 92–99.
- Watanabe, Y., Hedenquist, J.W., 2001. Mineralogic and stable isotope zonation at surface over the El Salvador Porphyry Copper Deposit, Chile. *Econ. Geol.* 96, 1775–1798.
- Williams, T.J., Candela, P.A., Piccoli, P.M., 1995. The partitioning of copper between silicate melts and two-phase aqueous fluids: an experimental investigation at 1 kbar, 800 °C and 0.5 kbar, 850 °C. *Contrib. Mineral. Petrol.* 121, 388–399.
- Wood, S.A., 1990a. The aqueous geochemistry of the rare-earth elements and yttrium: 1. Review of available low-temperature data for inorganic complexes and the inorganic REE speciation of natural waters. *Chem. Geol.* 82, 159–186.
- Wood, S.A., 1990b. The aqueous geochemistry of the rare-earth elements and yttrium: 2. Theoretical predictions of speciation in hydrothermal solutions to 350 °C at saturation water vapor pressure. *Chem. Geol.* 88, 99–125.
- Zimelman, D.R., Rye, R.O., Breit, G., 2005. Origin of secondary sulfate minerals on active andesitic stratovolcanoes. *Chem. Geol.* 215, 37–60. (this volume).