Barite-celestine intergrowths in Archean plutons: The product of oxidizing hydrothermal activity related to alkaline intrusions

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Abstract

Fine sulfate grains are disseminated within feldspar porphyries of the late Archean Kirkland Lake alkaline igneous complex in the Abitibi greenstone belt. Some grains are completely enclosed within hydrothermal biotite. Bulk compositions of the sulfates are intermediate in the BaSO₄-SrSO₄ series, the majority containing 15 to 20 mol% BaSO₄. Optical properties and the result of acid-etching indicate that they are mechanical mixtures of fine barite and celestine. The consistent bulk chemical compositions, δ^{34} S values (~+6‰), and high SO₄²⁻ contents in associated apatite indicate that they were formed as a barian celestine solid solution during oxidizing hydrothermal activity.

Lack of any major thermal events in post-Archean time and low ⁸⁷Sr/⁸⁶Sr values of sulfates indicate the formation of sulfates in late Archean time. Low metamorphic grade of the area and the occurrence of sulfates within biotites suggest that the sulfate formation was related to the intrusion of the alkaline complex.

INTRODUCTION

Near-surface environments in the Archean are considered to be generally reduced (e.g., Veizer et al., 1982; Walker et al., 1983). Archean sulfate is therefore believed to have been concentrated in (semi-) closed basins by biological oxidation of volcanic sulfide (e.g., Lambert et al., 1978; Reimer, 1980). Findings of barite in many Archean gold veins hosted dominantly by igneous rocks led to the recognition of Archean sulfate of nonbiological origin (Hattori and Cameron, 1986). This paper reports the occurrence of disseminated sulfates in late Archean intrusions at Kirkland Lake in the Abitibi greenstone belt and discusses their origin.

GEOLOGY

The Kirkland Lake area in the Abitibi greenstone belt represents one of the shallowest sections of Archean crust. Clasts of Timiskaming conglomerates, which host the intrusive complex, display evidence of prehnite-pumpellyite facies of regional metamorphism (e.g., Jolly, 1974). The intrusive complex has four major units: augite syenite, felsic syenite, feldspar porphyry, and quartz-feldspar porphyry (Fig. 1). U-Pb zircon ages from a porphyry sample indicate a maximum age of 2680 Ma (T. E. Krogh, personal communication, 1987). The complex is cut by volumetrically minor minette and vogesite dikes, and titanite from the minette dyke provides a minimum age of 2674 \pm 2 Ma (Wyman and Kerrich, 1987).

The reader may refer to Jensen and Langford (1985) and Hodgson (1983) for a discussion of regional geology and to Thomson (1950), Ploeger and Crocket (1982), and Hicks and Hattori (1988) for the geology of the mine area and the description of the gold mineralization.

Petrology of the feldspar porphyry unit

Monzodiorite porphyry is the predominant phase in the porphyry unit, and it contains ~25 vol% phenocrysts of (in order of decreasing abundance) plagioclase (3 mm), biotite (1.5 mm), green hornblende (<1 mm), and lesser amounts of magnetite (<0.1 mm), titanite (<0.4 mm), and apatite (<0.2 mm) microphenocrysts. Monzonite porphyry contains ~35 vol% phenocrysts of plagioclase (~3 mm), K-feldspar (~8 mm), biotite (1.5 mm), and minor amounts of green hornblende (1 mm) as well as magnetite (0.1 mm), titanite (<0.3 mm), and apatite (0.2 mm) microphenocrysts. The groundmass of both phases is composed mainly of albite, K-feldspar, and quartz with minor apatite and biotite. Chemical compositions of typical porphyry intrusions are listed in Table 1.

The porphyries, which host most of the gold ore in the camp, have undergone extensive alteration. However, rocks far from gold veins (>100 m), obtained mainly from the drill cores, are olive black (GSA color code: 5Y2/1) to dark grey (N3) and exhibit preserved minerals and textures that predate the alteration related to the gold mineralization. Foliation, veinlets, and fractures are virtually absent from the samples. Only the least-altered rock samples contain the sulfates described in this report.

Alteration observed in the least-altered samples includes formation of hematite and titanite on rims of magnetite, formation of fine-grained K-feldspar and albite on the margin of feldspar grains, partial recrystallization of groundmass feldspars, dissemination of minor ankerite in groundmass and the formation of secondary biotite (Hicks and Hattori, 1988).

Secondary biotite commonly replaces hornblende and primary biotite and overgrows the primary biotite. It is



Fig. 1. Geologic map of the Kirkland Lake area. The studied samples were collected from the area indicated in the inserted map. Modified after Ploeger and Crocket (1982).

characterized by its light color and high Mg ($X_{phl}^{biot} = \sim 70$ mol%; Fig. 2). Chemical compositions of secondary biotite vary within a section, and coarse one in the groundmass is greenish and rich in F (<1.8 wt%) and BaO (<2.8 wt%; Table 2).

OCCURRENCE OF SULFATES

Sulfates are found as common accessory minerals in various phases of the porphyry collected in this study (Fig. 1). Sulfates occur disseminated within secondary biotite (Figs. 3 and 4), in the margins of albite and K-feld-spar phenocrysts, and within the groundmass. Sulfate grains in biotite are commonly rectangular (1:2; Fig. 3), and some are elongated with aspect ratios up to 10:1 along the cleavage planes of biotite (Fig. 4). Grains in the groundmass are mostly anhedral and range in size from less than 2 to 20 μ m, but a single grain of ~100 μ m in length has been observed. Some grains in the groundmass enclose one or two minute grains of apatite, albite, and K-feldspar. An apatite inclusion in sulfate contains 0.9 wt% SO₃ (Table 3), which is higher than the SO₃ contents

in microphenocryst apatite. High SO_3 contents in apatite (Table 3) indicate an SO_4 -rich fluid during formation.

Sulfate grains are transparent under the microscope and have relatively high refractive indices and very low birefringence, $\delta < 0.006$, similar to those of apatite. There is no cleavage observed in the grains, although cleavage of {001} would be expected in a sulfate (Deer et al., 1963). They also show incomplete or undulatory extinction (Fig. 3c). Elongated sulfate grains in biotite do not show parallel extinction (Fig. 3b). The extinction under crossed polarizers is attained at an oblique angle (20° to 27°; Fig. 3c). These optical properties are not consistent with those of a single sulfate phase.

Chemical composition of sulfates

More than 40 analyzed grains of sulfate show only peaks of Ba, Sr, and S in the energy-dispersive spectra. Other cations, including Ca and Pb, are below detection limits (<0.3 wt%). Semiquantitative analyses show that most grains have similar chemical compositions; Ba/(Ba + Sr) molar ratios all fall in the range of ~0.15 to 0.20 (Table

Sample number	KDH83	KH414-12	KDH135
SiO ₂ (wt%)	64.12	61.41	62.27
TiO ₂ (wt%)	0.32	0.41	0.39
Al ₂ O ₃ (wt%)	14.55	14.22	14.64
FeO* (wt%)	1.72	2.40	1.94
Fe ₂ O ₃ (wt%)	1.44	1.47	1.95
MgO (wt%)	1.70	3.08	2.01
CaO (wt%)	2.70	3.33	3.28
Na ₂ O (wt%)	5.22	4.38	5.66
K₂O (wt%)	2.90	3.38	2.49
P ₂ O ₅ (wt%)	0.21	0.36	0.27
MnO (wt%)	0.06	0.07	0.07
S (wt%)	0.11	0.07	0.16
Ba** (ppm)	3531	3047	3169
Cr (ppm)	116	167	103
Zr (ppm)	260	217	256
Sr (ppm)	5666	2334	5934
Rb** (ppm)	73	115	66
Y (ppm)	16	17	14
Zn (ppm)	59	61	52
Ni (ppm)	14	58	16
V (ppm)	45	64	72
Sc** (ppm)	6.8	9.1	8.8
Cs** (ppm)	2.8	9.3	2.8
La** (ppm)	36	52	45
Ce** (ppm)	75	110	95
Nd** (ppm)	36	52	45
Sm** (ppm)	6.4	8.4	7.8
Eu** (ppm)	1.57	1.99	1.87
Tb** (ppm)	0.61	0.74	0.79
Dy** (ppm)	3.0	3.8	3.6
Ho** (ppm)	0.66	0.78	0.84
Yb** (ppm)	1.54	1.60	1.62
Lu** (ppm)	0.20	0.23	0.23
Total (wt%)	96.15	95.58	96.29

 TABLE 1.
 Chemical compositions of typical porphyry intrusions containing barium and strontium sulfates

Note: KDH83—Biotite monzodiorite porphyry, DDH 24 from 800-ft (~244-m) level at the Lake Shore property, 143–145 ft (~44 m) deep. KH414-12—Biotite monzodiorite porphyry, DDH 15 from 1500-ft (~457-m) level at the Lake Shore mine, 227 ft (~69 m) deep. KDH135—Hornblende-biotite monzodiorite porphyry, DDH 68 from 1000-ft (~300-m) level at the Lake Shore mine, 58 ft (~18 m) deep.

* Determined by titration technique.

** Determined by INAA at Ecole Polytechnique. All elements were determined by XRF, except as noted.

4; X-ray maps of $SrL\alpha$ and $BaL\alpha$ in Figs. 4b, 4c, 5a, and 5b).

Intragrain compositional variations were examined by electron microprobe and SEM. Backscattered-electron images and X-ray maps of SrL α and BaL α at 3000× to 4000× magnification (Figs. 4b, 4c, 5a, and 5b) do not reveal exsolution textures nor apparent inhomogeneity. Other grains, however, show patterns with slightly brighter and darker spots in backscattered-electron images corresponding to Ba- and Sr-rich domains (Table 4). Because celestine is far more soluble than barite at room temperature, samples were acid-etched (1:5 HCl) and specimens were examined by SEM; the solution was examined for its Sr and Ba contents by flame AAS. In two specimens, pure tabular barite was found in cavities that were once filled with sulfates (Fig. 6b). In most cases, blades of sulfates containing Ba and Sr remain in the sites (Fig. 6a), but the solution contained predominantly Sr. Further etching of the samples removed sulfates completely. The results indicate that some sulfate grains are made up of coarser



Fig. 2. Triangular diagram showing the composition of biotite, in terms of molecular FeO, MgO, and TiO_2 ; total Fe expressed as FeO. Solid circles = primary biotite; crosses = brown secondary biotite; open circles = green secondary biotite. Striped area = field of the compositions of primary biotite; dotted area = field of the compositions of secondary biotite.

barite and celestine ($\sim 1 \ \mu m$), but most are mixtures of much finer grains of the two sulfates. Because celestine is the major sulfate in the mixture, dissolution of celestine caused disintegration of grains and removal of sulfates from the sample.

Isotopic compositions of sulfates

S-isotope values of the sulfate are in a narrow range from +6 to +7‰ (Table 5b). The values are much larger than the values of sulfides in veins in the area; δ^{34} S values of sulfides in the auriferous veins range from -16 to -6‰ (Cameron and Hattori, 1987, and Table 5b), and the values in disseminated pyrite near the veins also are similarly low (Table 5b).

Sr was extracted by cold NaCl solution and the Srisotope compositions were determined. The values are low (0.7027 and 0.7040; Table 6).

DISCUSSION

There are three possible modes of origin of the sulfates in the samples: (1) a primary magmatic origin in which the sulfates crystallized from the melt, (2) a magmatichydrothermal origin, or (3) a supergene origin.

S-isotope data do not support the possibility of a supergene origin by oxidation of sulfides. Such sulfate would have low δ^{34} S values similar to the source sulfide, as sulfides in the area have low δ^{34} S (Table 5b). Instead, the observed values of δ^{34} S are similar to those for magmatic anhydrite (+6.5 to 9.5%; Arculus et al., 1983; Rye et al., 1984) and those for early-formed sulfate in the biotite-alteration zone in porphyry copper deposits, (+7.3 to 9.7%; Field and Gustafson, 1976; +6.6 to +8.1%; Shelton and Rye, 1982).

14100		
	RT31-F	880427-10
SiO₂ TiO₂	37.14 3.43	38.89 1.18
Al ₂ O ₂	13.87	14.27
Cr ₂ O ₃	n.d.	0.39
MgO	15.36	17.88
MnO	0.13	n.d.
FeO*	14.67	14.92
Na ₂ O	0.13	0.06
K₂O	8.96	9.35
CaO	n.d.	n.d.
BaO	1.38	0.21
F	n.d.	1.58
	n.d.	n.d.
	0.00	0.67
Total	95.07	98.06
Cations calculated Si ⁴⁺ Al ³⁺	on the basis of 11 ox 2.806 1.194	ygen equivalents 2.796 1.204
T site	4.000	4.000
Al ³⁺ Mg ²⁺ Mn ²⁺ Fe ^{2+*} Cr ³⁺ Ti ⁴⁺ M site	0.041 1.730 0.008 0.927 0.000 0.195 2.901	0.005 1.917 0.000 0.897 0.022 0.064 2.895
Na⁺ K⁺ Ca²⁺ Ba²⁺ A site	0.019 0.863 0.000 0.041 0.927	0.008 0.858 0.000 0.006 0.872
F ⁻ Cl O²-	0.000 0.000 11.000	0.359 0.000 10.641

TABLE 2. Compositions of secondary biotite associated with sulfates

Note: n.d., below the detection limit (<0.02 wt%).

Consistent bulk chemical compositions of the sulfates in the series $BaSO_4$ -SrSO₄ also support this view. Because there is immiscibility in the series at temperatures below 100 °C (Malinin and Urusov, 1983), sulfates formed in supergene conditions would not have intermediate compositions. Their occurrence within biotite and the lack of sulfate veinlets in the samples are also consistent with their formation at high temperatures.

Sulfates may be formed as a primary phase in magmas whose f_{02} is near the NNO buffer (Carroll and Rutherford,

TABLE 3. Chemical compositions of apatite (in wt%)

	A	в	С
CaO	54.74	54.20	55.91
FeO	n.d.	n.d.	
SrO	n.d.	n.d.	n.d.
P ₂ O ₅	42.04	41.35	41.61
SO3	0.24	0.39	0.90
CI	n.d.	n.d.	
Total	97.02	95.94	98,42

Note: n.d., not detected; —, not determined. (A) Apatite microphenocrysts in sample KDH136 (DDH 93 from 1000-ft (\sim 300-m) level, 6–7 ft (\sim 2 m) deep at the Lake Shore property). (B) Apatite microphenocrysts in sample KH414-12. (C) Apatite enclosed in the sulfate grains in groundmass.



Fig. 3. Photomicrograph showing typical occurrence of sulfates within biotite (sample no. KDH83). (a) Plane-polarized light; (b) and (c) cross-polarized light. Scale bar is equivalent to 100 μ m. Note that the sulfate grains (marked S) do not show parallel extinction (b) and their extinction is incomplete (c).

1987). The magmas from which the Kirkland Lake porphyries were formed were apparently oxidized, as indicated by early formation of titanite and magnetite together with hornblende (Noyes et al., 1983) and high Mg/Fe values in the primary biotite (Fig. 2; Chivas, 1981). However, the occurrence of sulfates in secondary biotites and the variation in Sr in the porphyry samples, >4000 ppm in sulfate-bearing samples (Table 1) and ~1000 ppm in



TABLE 4. Chemical compositions of sulfates (electron-microprobe analysis)

	A	в	С	D
SrO (wt%)	41.9	45.0	43.7	50.9
BaO (wt%)	16.9	13.3	14.8	6.4
CaO (wt%)			0.1	0.3
SO3 (wt%)	41.2	41.7	41.5	42.7
SrSO ₄ (mol%)	78.6	83.35	81.14	91.30
BaSO, (mol%)	21.4	16.65	18.59	7.77
CaSO ₄ (mol%)			0.26	0.96

Note: (A) Grains within secondary biotite (sample KDH83). (B) Darkest spot of the grain that showed inhomogeneity in backscattered-electron image (sample KDH84). (C) Brightest spot of the same grain as B (sample KDH84). (D) Sr-rich grain in the groundmass (sample KDH135).

sulfate-free samples, suggest a hydrothermal origin of the sulfates.

Biotite is a common metamorphic mineral. However, the biotite in the Kirkland Lake intrusive complex is not a product of regional metamorphism because of low regional metamorphic grade (e.g., Jolly, 1974). The formation of sulfates in the intrusions is, therefore, related to the emplacement of intrusive complex. Low ⁸⁷Sr/⁸⁶Sr ratios in the sulfates support this view. Because the porphyry unit contains a significant amount of Rb (70 to 300 ppm), any sulfate formed in post-Archean time would be rich in radiogenic ⁸⁷Sr.

Decompression of magmas during their ascent allows a release of volatiles. SO_2 is the principal S-bearing species in the volatile phase of oxidized magmas, and it forms SO_4^2 through a hydrolysis reaction. The release of volatiles from the studied porphyries is supported by the presence of miarolitic cavities (Hicks and Hattori, 1988) and titanization and hematitization along cracks and margins of magnetite microphenocrysts.

Sulfates display retrograde solubilities at various temperature ranges. Solubilities of celestine are retrograde at temperatures below ~400 °C (Strubel, 1966). Solubilities of barite in dilute solutions, <1m NaCl, are retrograde at temperatures higher than 100 °C (Blount, 1977). Retrograde solubilities of barite are also reported in saline fluids at temperatures between 350 and 450 °C (Strubel, 1967). The occurrence of the mixture of celestine and barite in the groundmass of the studied samples may suggest that the pervasive hydrothermal activity ceased at temperatures higher than 400 °C. Lack of retrograde alteration of biotites may support this interpretation.

Anhydrite is the most common sulfate in igneous rocks either as phenocrysts or alteration product. The lack of anhydrite in studied samples may be attributed to low Ca/Sr and Ca/Ba ratios of the host intrusions (Table 1) and high CO_2 of fluid responsible for the formation. Solubilities of anhydrite are over an order of magnitude larger than those of celestine and barite in a solution of a given salinity (Strubel, 1966, 1967; Blount and Dickson, 1969), which suggests that precipitation of anhydrite requires approximately the concentration of Ca^{2+} to be ten

Fig. 4. (a) Backscattered-electron image, (b) $SrL\alpha$ X-ray map, and (c) $BaL\alpha$ X-ray map of sulfate grains within biotite shown in Fig. 3.



Fig. 5. (a) $SrL\alpha$ X-ray map and (b) $BaL\alpha$ X-ray map of sulfate in groundmass. Q = quartz, K = K-feldspar.

times larger than Sr^{2+} . The host porphyry rocks are rich in Sr and Ba, but low in Ca (Ca/Sr molar ratios, 7 to 8). Furthermore, K-feldspar in the Kirkland Lake intrusive complex is rich in BaO (<3.01 wt%) and SrO (<2.88 wt%). Recrystallization of feldspars during hydrothermal activity would have released Ba²⁺ and Sr²⁺ to the fluids.

TABLE 5a. S-isotope compositions of sulfates

Sample	Total S (wt%)	$\delta^{34}S_{CDT}$ (‰)
KDH82	0.11	+6.2
KDH119	0.08	+6.2
KDH130	0.09	+6.7
KH414-12	0.07	+7.0
RT32	0.12	+6.0
		+6.1*



Fig. 6. Scanning-electron-microscope image of sulfate after etching with acid. (a) Tabular barite crystals (s) in biotite (bt) (sample RT-39). (b) Sulfate blades (s) containing Sr and Ba in biotite (bt). Photomicrographs of the grain shown in Figs. 3 and 4 after acid etching.

Finally, fluids were rich in CO_2 , indicated by minor but ubiquitous occurrences of ankerite in the groundmass. Ca^{2+} concentration of such fluids would have been buffered by carbonate, which would have prevented the formation of anhydrite.

TABLE 5b. S-isotope (compositions o	f sulfides
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Sample	Description	δ ³⁴ S _{CDT} (‰)
KH85022-04	North vein	-11.7
KH860220-31	Syenite, 0.5 m from vein	-10.8
KH850220-60	Network vein in porphyry	-3.3
KH850220-71	Dissemination of pyrite in porphyry	-9.6
KH860529-09	Pyrite in porphyry, 0.3 m from South vein	-9.4
KH860529-10	Pyrite in South vein	-10.4
KH880427-16	Molybdenite-rich part of the main vein	-9.1
KH880427-15	Pyrite cubes in fault gouge near the North vein	-6.7
RT-05	Sericitized-pyritized wallrock	-10.3
RT-12	Hematitized wallrock	-2.8

Sample	⁸⁷ Sr/ ⁸⁶ Sr
KDH83	0.70271 ± 0.00002
KDH119	0.70403 ± 0.00006

TABLE 6. Sr-isotope compositions of sulfate

The occurrence of the primary sulfate minerals has been recognized within only historic lavas (e.g., Arculus et al., 1983; Luhr et al., 1984). The absence of anhydrite phenocrysts in prehistoric igneous rocks is attributed to weathering because of the high solubility of sulfate (Arculus et al., 1983). Similarly, magmatic-hydrothermal sulfates are common in Phanerozoic intrusions, but their occurrence in Archean terranes is very limited. This is partly due to erosion of the upper section of Archean crust and partly due to the retrograde solubilities of many sulfates. Most magmatic and magmatic-hydrothermal sulfates would have been dissolved during the waning stage and/or later hydrothermal activity.

In Phanerozoic porphyry deposits, dissolution of earlier formed magmatic-hydrothermal sulfates affected the chemistry of subsequent hydrothermal fluids and resulted in redistribution of metallic minerals (e.g., Shelton and Rye, 1982). Under the generally reducing Archean conditions, the dissolution of sulfate would have had a profound effect on the chemistry of subsequent hydrothermal fluids at <400 °C by raising their f_{02} to that of the H₂S-SO₄²⁻ buffer. Increase in oxidation of fluids to the H₂S-SO₄²⁻ buffer during the waning stage of hydrothermal activity would result in transport of only selected elements, because most metals are less soluble in more oxidized fluids and at lower temperatures. Au and Mo belong to this group of elements that could be concentrated by oxidized fluids.

SUMMARY

1. Physically mixed grains of barite-celestine occur disseminated in the least-altered monzonitic porphyry intrusions of Archean age at Kirkland Lake, Ontario.

2. The occurrence of sulfates within secondary biotite, bulk Ba/Sr ratios and δ^{34} S and 87 Sr/ 86 Sr ratios of the sulfates indicate that they were most likely formed at high temperatures, >400 °C, as an intermediate-composition member in the BaSO₄-SrSO₄ series during the emplacement of the alkaline complex in late Archean time.

Many epizonal intrusions of Archean age should be carefully studied to identify the occurrence of sulfate as it may be more common than is generally realized.

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APPENDIX 1. ANALYTICAL METHODS

Major- and trace-element contents of whole-rock samples were determined by xRF on samples fused with a Li_2CO_3 - $Li_2B_4O_7$ mixture. Fe²⁺ contents were determined by a standard titration technique. Chemical analyses of minerals were carried out using a JEOL-733 electron microprobe equipped with a Tracor Northern 5500 energy-dispersive spectrometer and Tracor Northern 5600

automation. For all samples, the general operating conditions were as follows: 15-kV accelerating voltage, 25-nA beam current, 50-s count time (or 0.5% precision, whichever was achieved first). For the apatites and biotites, data reduction was carried out on data collected with wavelength spectrometers using the Tracor-Northern 5600 program TASK; standards were apatite (P, Ca) and anhydrite (S) for apatites, chlorite (Al, Si, Mg), biotite (Fe, Ti, K), diopside (Ca), chromite (Cr), albite (Na), willemite (Mn), tugtupite (Cl), sanbornite (Ba), and fluorellestadite (F) for biotite analysis. In the case of sulfates, compositions could only be determined semiguantitatively because of the small grain size (generally less than 10 μ m) and the decomposition of the samples under high beam current. For this purpose, the semiquantitative program sso in the Tracor-Northern 5500 was used for the analysis of the energy-dispersive spectra of the minerals. Ca contents were calculated using wavelength-spectrometer data in conjunction with the TASK program. SEM study was carried out using SEMCO Nanolab 7 attached KEVEX energy-dispersive X-ray analyzer. The accelerating voltage of the beam was set at 15-kV throughout the analysis.

For isotope analysis, S in the samples was extracted by "Kibareagent" (Sasaki et al., 1979). Because no sulfides were observed in sections under microscope, the values of total S are considered to be those for sulfates. To ensure the result, sulfate S was leached from several samples by 5% HCl solution and then precipitated as BaSO₄. SO₂ for isotope analysis was extracted from the BaSO₄ by mixing with Cu₂O and SiO₂ and combusting the mixture at 1000 °C. The isotopic compositions were obtained using VG micromass 602E, and the data are expressed with respect to the Canon Diablo meteorite standard using the familiar δ notation. Overall analytical precision is $\pm 0.1\%$.

Sr was extracted from the rock powder (less than 2 μ m) by constantly stirred cold 2N NaCl solution for 2 d, which prevented dissolution of Sr from carbonates and silicate minerals. Sr was then separated from the NaCl solution using a cationexchange resin and then analyzed by mass spectrometry on a Finnigan Mat 261.