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OLDHAMITE IN ENSTATITE ACHONDRITES (AUBRITES)

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Abstract: Properties of oldhamite (ideally CaS) in aubrites are summarized and compared to oldhamite in enstatite chondrites. The origin of the high REE abundances in aubritic oldhamite and the diversity of REE abundance patterns is addressed. Low CaS/silicate partition coefficients indicate that oldhamite in enstatite achondrites cannot have gained its high REE concentrations during igneous differentiation processes. However, the observed REE abundance patterns in oldhamite can be explained by REE condensation into CaS grains in the solar nebula. The very high melting point of oldhamite plausibly led to its preservation and prevented major exchange reactions of the oldhamite with other aubritic minerals during the short differentiation and metamorphism period on the aubrite parent body. Thus, oldhamite in aubrites is probably a slightly metamorphosed relict condensate, as suggested earlier.

1. Introduction

The mineralogy and chemistry of the highly reduced enstatite chondrites and enstatite achondrites (aubrites) are distinctive compared to that of other meteorite types and terrestrial rocks (MASON, 1966; KEIL, 1968). Table 1 lists some of the minerals observed in the primitive EH and EL chondrites as well as in aubrites, which are silicate rocks from a differentiated parent body. The occurrence of some of the exotic sulfides and metallic phases is explained by the formation of these meteorites under highly reducing conditions. Highly reducing conditions imply that oxygen was depleted and lithophile elements, which are normally observed in oxide and silicate phases, occur (partially) as sulfides (Fe, Mg, Mn, Ca, Ti, REE), as nitrides (Si, Ti), or alloyed with metallic FeNi (*e.g.*, Si).

Among the sulfide phases, oldhamite (ideal formula CaS) is of particular interest for two reasons. First, pure CaS is very refractory with a melting point of 2450°C. It is calculated (LARIMER, 1975; LARIMER and BARTHOLOMAY, 1979; LODDERS and FEGLEY, 1993) to be among the first condensates in a reduced solar gas (for C/O>1, CaS condenses at 1379 K at 10⁻³ bars total pressure). Thus, oldhamite is analogous to hibonite (CaAl₁₂O₁₉) and perovskite (CaTiO₃) the two most refractory Ca-bearing phases in Ca,Al-rich inclusions in carbonaceous chondrites (*e.g.*, KORNACKI and FEGLEY, 1984, 1986).

Second, oldhamite is an important host phase for the rare earth elements (REE) in enstatite chondrites and achondrites. Typically, REE abundances in oldhamite are about 100 times CI-chondritic and a variety of REE abundance patterns are observed (*e.g.*, LARIMER and GANAPATHY, 1987; FLOSS *et al.*, 1990; LODDERS and PALME 1991; KURAT *et al.*, 1992; FLOSS and CROZAZ, 1993; LODDERS *et al.*, 1993; WHEELLOCK *et al.*, 1994; CROZAZ

Table 1. Minerals in enstatite chondrites and achondrites.

Mineral name	Chemical formula	Abundances (wt%)		
		EH3/4	EL5/6	Aubrites
Orthoenstatite	Mg ₂ Si ₂ O ₆	46–52	62–67	77–97
Clinoenstatite	Mg ₂ Si ₂ O ₆	trace	n.r.	trace
Diopside	MgCaSi ₂ O ₆	trace	n.r.	0.2–7.8
Olivine	Mg ₂ SiO ₄	trace	n.r.	0.3–9.8
Plagioclase	>Ab90	1.5–16	6.4–9.2	0.3–14
Roedderite	(Na,K) ₂ Mg ₅ Si ₁₂ O ₃₀	trace	n.r.	trace
Quartz	SiO ₂	trace	trace	trace
Cristobalite	SiO ₂	1.5–4	n.r.	trace
Tridymite	SiO ₂	trace	trace	n.r.
Fluor-Richterite	Na ₂ CaMg ₅ Si ₈ O ₂₂ (F,OH) ₂	trace	n.r.	trace
Alabandite	(Mn, Fe)S	n.r.	0.25–0.4	0.25–0.4
Caswellsilverite	NaCrS ₂	trace	n.r.	trace
Daubreelite	FeCr ₂ S ₄	0–0.2	n.r.	trace
Djerfisherite	K ₃ (Cu,Na)(Fe,Ni) ₁₂ (S,Cl) ₁₄	trace	n.r.	trace
Niningerite	(Mg,Fe)S	0.2–11.2	n.r.	n.r.
Oldhamite	CaS	<0.1–2	0.07–0.9	0–0.4
Sphalerite	ZnS	trace	trace	trace
Troilite	FeS	5.8–13	4.6–9.5	0.4–1.5
Heideite	FeTi ₂ S ₄	n.r.	n.r.	trace
Cohenite	(Fe,Ni) ₃ C	trace	trace	n.r.
Kamacite	α-FeNi	17.5–23.8	13.3–24.2	0–2.47
Perryite	(Ni,Fe) ₅ (Si,P) ₂	trace	trace	trace
Schreibersite	(Fe,Ni) ₃ P	0.1–1	0.66–1	trace
Taenite	γ-FeNi	trace	trace	trace
Graphite	C	0.03–0.6	0.1–0.4	trace
Lawrencite	(Fe,Ni)Cl ₂	trace	n.r.	n.r.
Nierite	Si ₃ N ₄	trace	n.r.	n.r.
Osbornite	TiN	trace	trace	trace
Sinoite	Si ₂ N ₂ O	n.r.	0–0.14	n.r.

trace: less than 0.1 wt%. n.r.: not reported. Data are compiled from KEIL (1968), WATTERS and PRINZ (1979) as well as many other sources.

and LUNDBERG, 1995). The high REE abundances and different REE abundance patterns in oldhamite in enstatite chondrites can be plausibly explained by condensation processes in a reduced solar gas (LARIMER *et al.*, 1984; LODDERS and FEGLEY, 1993). However, the origin of oldhamite in aubrites and the origin of its high REE content is still under debate (see below).

The aubrite parent body underwent igneous differentiation after metal and sulfides segregated from the silicates to form a core. The large crystal size of enstatite in some aubrites and the igneous texture of silicate assemblages indicate that some crystallization process took place on the aubrite parent body (*e.g.*, OKADA *et al.*, 1988). In order to model crystallization processes the REE distribution in aubritic minerals and the partition coefficients for the relevant mineral/melt systems need to be known.

Several measurements of REE abundances in bulk aubrites and in their individual mineral phases are available (*e.g.*, SCHMITT *et al.*, 1963; MASUDA, 1967; WOLF *et al.*,

1983; GRAHAM and HENDERSON, 1985; FLOSS and CROZAZ 1993; LODDERS *et al.*, 1993; WHEELOCK *et al.*, 1994). Likewise, the REE mineral/melt partition coefficients for pyroxene, olivine, diopside, and plagioclase are available (KENNEDY *et al.*, 1993; GRUTZECK *et al.*, 1974; DRAKE and WEILL, 1975). For modeling the igneous differentiation history of the aubrite parent body (APB) CaS/silicate partition coefficients are also needed, because the REE are strongly concentrated in oldhamite. Under highly reducing conditions the REE may not only partition between CaS and silicate but also between FeS and silicate. Some REE (and in particular Eu) may have been removed by an Fe-FeS melt during formation of the metal-sulfide core (LODDERS and PALME, 1989, 1990), which could explain the frequently observed negative Eu-anomaly in bulk aubrite samples (*e.g.*, WOLF *et al.*, 1983). To model this potential fractionation, REE FeS/silicate partition coefficients are also required.

The origin of oldhamite and its role during igneous differentiation is still unclear. Two different schools of thought exist on the origin of oldhamite in aubrites. One is that oldhamite crystallized from a melt during igneous differentiation on the aubrite parent body. Another is that oldhamite is a relict condensate, altered to a lesser or greater extent during igneous activity on the APB. The arguments pro and con for these opinions are summarized in the present paper. In addition, some properties of oldhamite and a model of the differentiation of the aubrite parent body are discussed.

2. Occurrence and Composition of Aubritic Oldhamite

2.1. Occurrence

In the brecciated aubrites, oldhamite mainly occurs as single grains in the matrix and is relatively easily identified by its pinkish color, or when weathered, greenish appearance. Most grains are $\sim 100 \mu\text{m}$ to mm sized, but large grains up to 2 cm are occasionally observed (WHEELOCK *et al.*, 1994). Other sulfides (*e.g.*, troilite, alabandite) can be associated with oldhamite. In rare cases, oldhamite occurs as inclusions in pyroxene (FLOSS and CROZAZ, 1993; KURAT *et al.*, 1992). It is interesting to note that most oldhamite occurs as individual grains and does not display eutectic textures when in association with other sulfides or metal, as might be expected if eutectic melting in the CaS-FeS-Fe system had occurred (VOGEL and HEUMANN, 1941).

The oldhamite often contains fine blebs of troilite and ferroan alabandite, which in turn may contain small inclusions or associations with caswellsilverite (LODDERS *et al.*, 1993; WHEELOCK *et al.*, 1994). The modal abundance of oldhamite is typically less than one percent and, because oldhamite easily decomposes by aqueous weathering, aubrite finds often do not contain any detectable oldhamite.

2.2. Major element composition of aubritic oldhamite and equilibration temperatures

The major element composition of oldhamite in aubrites is given by WATTERS and PRINZ (1979). Stoichiometric CaS contains 55.55 wt% Ca but oldhamite in aubrites contains only 52.0–54.8 wt% Ca because MnS (~ 1 wt% Mn) and MgS (~ 0.3 wt% Mg) are dissolved in CaS. The FeS content in oldhamite is generally very low (Fe < 0.02 wt%) in accord with CaS-FeS phase diagram studies by SKINNER and LUCE (1971), who found no detectable solid solution of FeS in CaS. A comparison of the Ca, Mg, and Mn content of

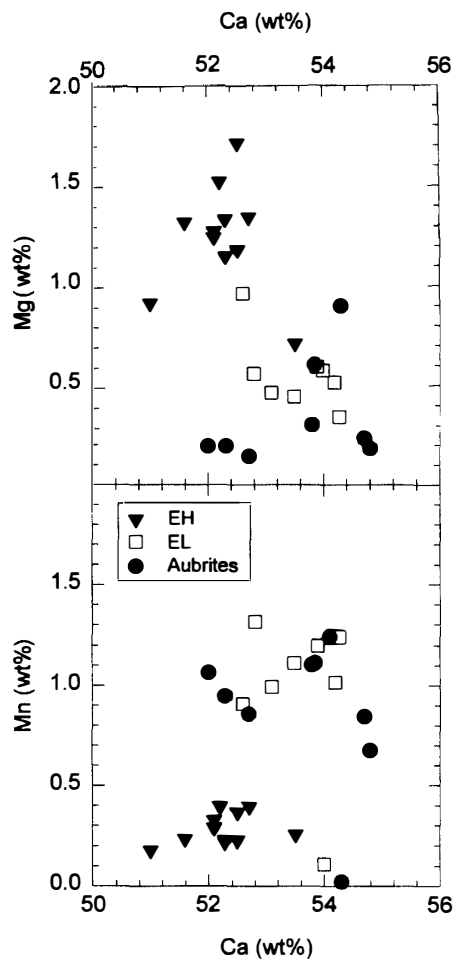


Fig. 1. Abundances of Mg, Mn, and Ca in oldhamite from enstatite chondrites and achondrites. The Mg content decreases and the Mn content increases from EH-chondrites to EL-chondrites to aubrites. Data from KEIL (1968), LARIMER and GANAPATHY (1987), LEITCH and SMITH (1982), WATERS and PRINZ (1979), OKADA et al. (1988), LODDERS et al. (1993).

aubritic oldhamite and oldhamite in enstatite chondrites is given in Fig. 1. LARIMER and GANAPATHY (1987) pointed out that the Mn and Mg contents of oldhamite in enstatite chondrites reflect the degree of metamorphism. With increasing degree of metamorphism (*i.e.*, increasing temperature), the Mn content increases and the Mg content decreases. The highest Mg content in oldhamite is found for the more primitive EH-chondrites, oldhamite in the more metamorphosed EL-chondrites has less Mg, and aubritic oldhamite is somewhat lower in Mg than EL-chondritic oldhamite. The inverse trend is observed for Mn in oldhamite, where EH-chondrites have the lowest Mn abundance and EL-chondrites and aubrites have higher and similar Mn contents.

One might expect that oldhamite of EL-chondrites and aubrites, which have experienced higher metamorphic temperatures than EH-chondrites, contains higher amounts of Mg, because the phase diagram studies by SKINNER and LUCE (1971) indicate a higher solubility of MgS in CaS at higher temperatures. However, in EL-chondrites and aubrites, niningerite ((Mg,Fe)S) is basically absent so that their study of MgS solubility in CaS for coexisting sulfides is not applicable. If niningerite were ever present in the EL-chondrite parent body or in the APB, the absence of niningerite in EL-chondrites and aubrites could be explained by oxidation of MgS by FeO or MnO from the silicates during metamorphic episodes.

SKINNER and LUCE (1971) developed the CaS-MnS-FeS geothermometer to determine the closure temperature of oldhamite in EL-chondrites. The similar compositions of EL-chondritic and aubritic oldhamite and the coexistence of small inclusions of troilite and alabandite in aubritic oldhamites allows us to apply the same geothermometer to aubrites (Fig. 2). Although REE distributions in aubritic minerals indicate that oldhamite is either partially or totally unequilibrated with respect to silicate phases (see below), we may assume that oldhamite is in local equilibrium with the associated sulfides and sulfide inclusions. The equilibration temperature can be calculated from:

$$T(^{\circ}\text{C})=603+13095\cdot X_{\text{MnS}}-119050\cdot X_{\text{MnS}}^2, \quad (1)$$

where X_{MnS} is the mole fraction of MnS in CaS coexisting with MnS (alabandite) and FeS (troilite). The presence of less than 0.5 wt% Mg in most aubritic oldhamites (see Fig. 1) may have some effect on the solvus but was neglected here.

The minimum equilibration temperatures for aubrites, calculated for the average oldhamite composition given by WATTERS and PRINZ (1979), are between 700–800°C. The exception is the Pena Blanca Spring aubrite, for which Watters and Prinz report

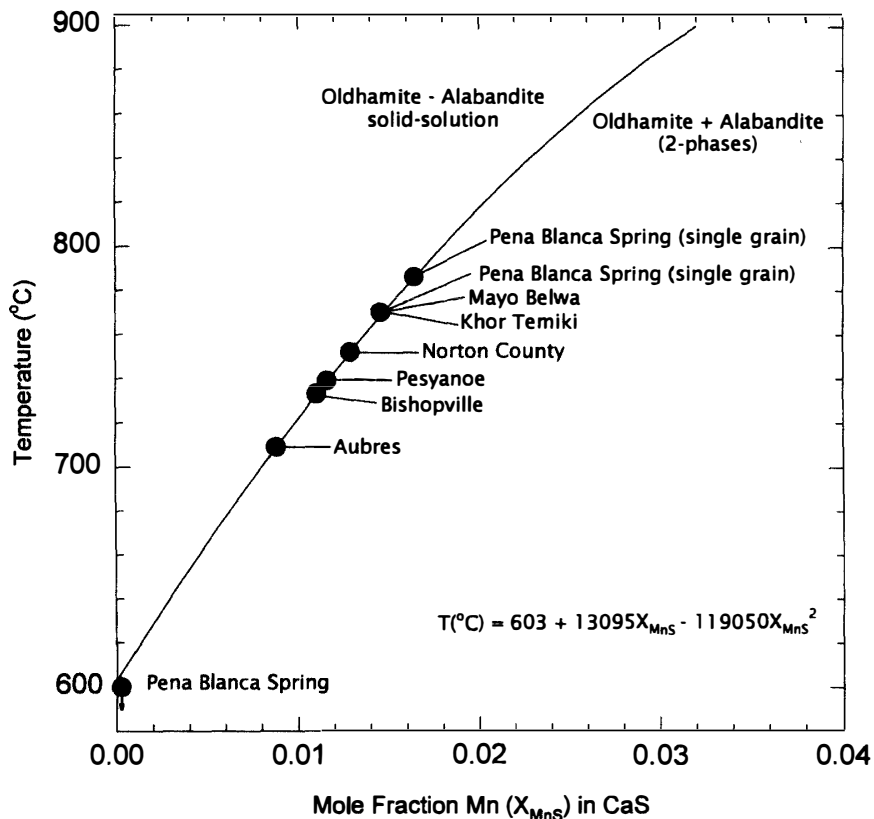


Fig. 2. Equilibration closure temperatures calculated with the CaS-MnS-FeS geothermometer of SKINNER and LUCE (1971). Data for aubritic oldhamite indicate equilibrium temperatures between 700 and 800°C. Data from WATTERS and PRINZ (1979), data for oldhamite single grains are from LODDERS *et al.* (1993).

<0.02 wt% Mn in oldhamite, which indicates closure of the CaS-MnS system below 600°C. However, the Mn content determined in two individual oldhamite grains from Pena Blanca Spring (LODDERS *et al.*, 1993) yield 770 and 790°C, which fall into the general trend. It would be interesting to determine the equilibration temperatures for oldhamite grains displaying different REE abundance patterns (see next paragraphs) but unfortunately to date, no published ion probe analyses of meteoritic oldhamites have included major elements (*e.g.*, Mg, Mn, Fe).

2.3. REE in oldhamite and other phases

The oldhamite from aubrites has been analyzed by several groups for trace elements, often with the main focus on the REE and actinides (MURRELL and BURNETT, 1982; BENJAMIN *et al.*, 1984; FLOSS *et al.*, 1990; KURAT *et al.*, 1992; FLOSS and CROZAZ, 1993; LODDERS *et al.*, 1993; WHEELOCK *et al.*, 1994). REE and actinides abundances are generally about 50–200 times CI-chondritic, which implies that these elements are almost completely located in oldhamite, even though the modal abundance of oldhamite is about one percent or less. FLOSS and CROZAZ (1993) report ten different REE abundance patterns in aubritic oldhamite, and different patterns were even observed in oldhamites from the same meteorite.

Other REE bearing phases are albitic plagioclase with light REE abundances that are about 0.01–1 times CI-chondritic and Eu about 5–10 times CI, and diopside, which in most cases has a bow shaped REE pattern at about 0.1–10 times chondritic and an Eu-depletion. However, other patterns for diopside were also found (FLOSS and CROZAZ, 1993).

The high REE abundances and the diversity of REE patterns in oldhamite make representative sampling of the brecciated aubrites critical for determining the whole rock REE patterns in aubrites. The variety of bulk REE abundance patterns described in the literature (*e.g.*, SCHMITT *et al.*, 1963; MASUDA, 1967; WOLF *et al.*, 1983) probably depends on how much oldhamite was present in each whole rock sample analyzed, on the REE abundance patterns in the oldhamites contained in the sample, and to some extent also on the amount of plagioclase present in the sample. An example of a mass-balance calculation to model bulk REE abundances is given by LODDERS *et al.* (1993) for the Pena Blanca Spring aubrite.

3. REE CaS/silicate and FeS/silicate Partition Coefficients: Experimental Data

As mentioned above, REE abundances in aubritic minerals and REE silicate/melt partition coefficients are both known. Other necessary data for modeling aubrite petrogenesis are CaS/melt and FeS/melt partition coefficients for the REE. These are discussed next. The experimental details are described elsewhere (LODDERS *et al.*, 1990; LODDERS, 1991, 1995) and here only a brief summary is given, together with a comparison of other experimental results from JONES and BOYNTON (1983) and DICKINSON *et al.* (1990a, b, 1991). FeS/silicate and CaS/silicate partition coefficients are shown in Figs. 3 and 4, and some experimental results are listed in Tables 2 and 3. The partition coefficients (*D*) are expressed as weight ratios of concentration in sulfide over concentration in silicate matrix.

Table 2. Experimental conditions, concentrations of REE in iron sulfide and silicate, and REE FeS/silicate partition coefficients (*D*) in runs containing Al-foil as an oxygen getter.

Run	EQ9 ^b			EQ10 ^b			EQ4			EQ3		
T(°C)	1200			1200			1200			1200		
Time(h)	5			5			2			2		
X _{FeO} (sil.) ^a	0.042			0.039			0.106			0.109		
log fO ₂	-14.7			-14.8			-12.0			-12.3		
log fS ₂	-5.8			-5.8			-4.4			-3.8		
	Fe-S	Sil.	D	Fe-S	Sil.	D	Fe-S	Sil.	D	Fe-S	Sil.	D
Fe wt%	62.33	3.68	16.9	61.0	3.59	17	66.11	9.57	6.9	64.89	9.73	6.7
Sc ppm	<0.2	25.6	<0.007	<0.085	24.2	<0.004	<0.12	20.7	<0.006	<0.09	21.9	<0.004
La ppm	41	140	0.29	74.1	156	0.48	0.28	195	0.0014	0.13	221	0.0006
Pr ppm	142	433	0.33	280	545	0.51	0.7	686	0.001	–	753	–
Nd ppm	80	326	0.25	205	536	0.38	<32	692	–	–	710	–
Sm ppm	4.7	27.9	0.17	8.8	43.3	0.20	0.028	38.2	0.0007	<0.012	34.6	<0.0004
Eu ppm	10.5	12.3	0.85	16.5	15.3	1.08	0.096	27.6	0.0035	0.12	40.4	0.003
Gd ppm	66	818	0.081	155	1080	0.14	<4	1120	<0.0036	–	1140	–
Tb ppm	10	162	0.062	21	222	0.094	<0.5	178	<0.0028	–	190	–
Dy ppm	2.6	44.7	0.058	3.1	37.8	0.082	<0.027	34	<0.0008	<0.03	40	<0.0007
Ho ppm	1.53	58.8	0.026	1.9	39.6	0.048	–	39	–	–	44.2	–
Yb ppm	5.7	175	0.033	5.7	181	0.032	<0.2	156	<0.0012	<0.14	161	<0.0009
Lu ppm	0.31	26.2	0.012	0.48	37.5	0.013	0.14	31.7	0.0044	<0.04	34.6	<0.001

INAA data (LODDERS, 1991). a: mole fraction FeO in silicate after the experiment. b: FeS exsolved from Si-bearing Fe-metal. – stands for not analyzed.

The REE FeS/silicate partition coefficients decrease from light to heavy REE, except for Eu, which partitions about ten times stronger into the sulfide than neighboring Sm or Gd (Fig. 3). This trend was found in all studies. However, JONES and BOYNTON (1983) report higher FeS/silicate partition coefficients than LODDERS (1991, 1995) or DICKINSON *et al.* (1990a, b, 1991). This may be due to different sulfide and silicate compositions or different redox conditions used in the experiments. Experiments conducted under more oxidizing conditions yield lower REE FeS/silicate partition coefficients than do reducing experiments where Al-foil was present as an oxygen getter (LODDERS, 1991, 1995). However, under reducing conditions appropriate to aubrite formation, the REE clearly develop chalcophile trends.

The determination of partitioning coefficients under reducing conditions is generally difficult. Two experimental approaches have been used to measure REE CaS/silicate partition coefficients. In the method used by JONES and BOYNTON (1983) and by DICKINSON *et al.* (1990a, b, 1991) a charge of REE-spiked silicates, FeS₂, and Mg or Al as a reducing agent was equilibrated in evacuated sealed silica tubes. In this system, a (Ca, Mg)S liquid is formed by sulfurization of the silicates. The resulting (Ca, Mg)S only contained up to 30 wt% Ca, which is much more Mg-rich than any oldhamite found in aubrites (Mg < 1 wt%). In the method employed by LODDERS (LODDERS *et al.*, 1990; LODDERS, 1991), a Ca-free but otherwise enstatite chondrite-like silicate, spiked with REE, was equilibrated with CaS in evacuated silica tubes. In these experiments, solid CaS equilibrates with liquid silicate and the resulting CaS (48–50 wt% Ca) contained only up to 5 wt%

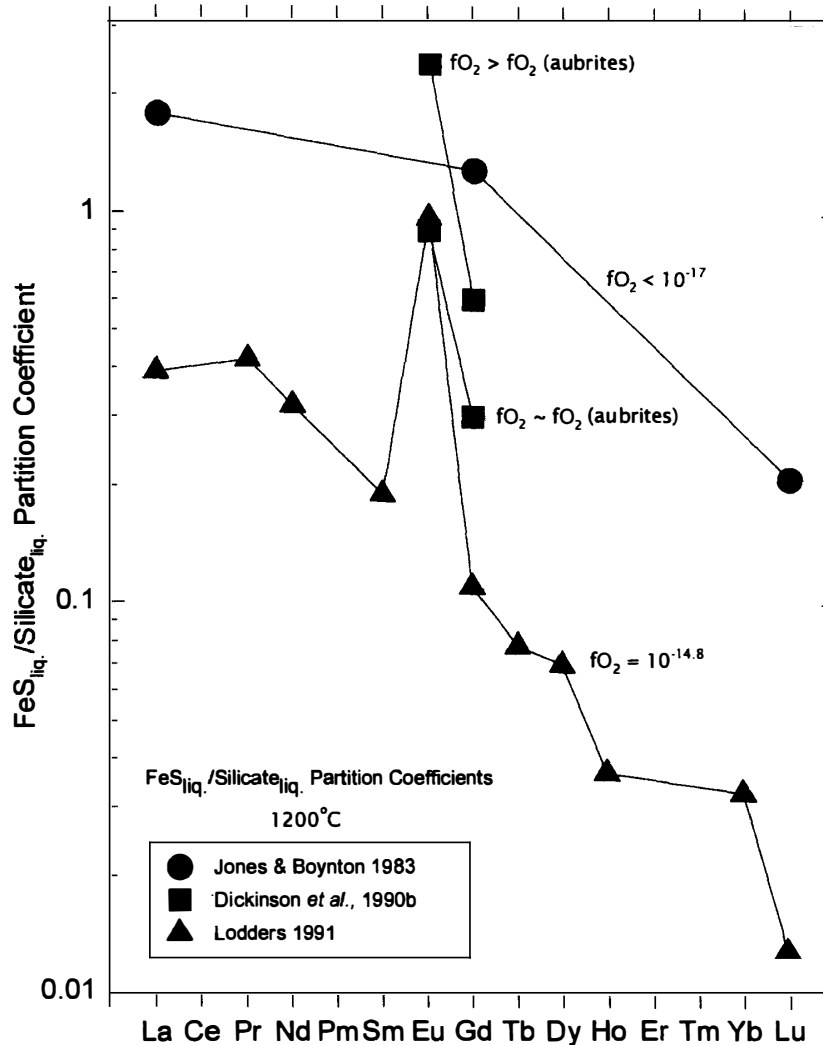


Fig. 3. *FeS/silicate partition coefficients for the REE at 1200°C under reducing conditions. The oxygen fugacities prevailing during the experiments are indicated in the graph. DICKINSON et al. (1990b) do not mention which fO_2 they assume for aubrite formation. Assuming that an fO_2 of 5 log units below the iron-wüstite buffer is representative for aubrite formation, then fO_2 (aubrites) $\sim 10^{-17}$ at 1200°C.*

Mg from exchange reactions with the silicate. However, the silicate matrix in the experiments by LODDERS *et al.* (1990) contained up to 15 wt% CaO after the experiment, thus making the silicate more calcic than bulk silicates observed in aubrites.

Since reaction times in silica tube experiments are limited by the stability of the quartz glass at reaction temperatures, it was necessary to check if equilibrium was obtained. Reversal experiments by LODDERS, using REE-spiked CaS and initially REE-free silicate, gave somewhat higher CaS/silicate partition coefficients (about a factor of 2 for Eu and ~ 4 for Sm and Gd, see Table 3). This indicates that equilibrium was not reached within the duration of the experiments. Thus, results of the reversal experiments represent upper limits for REE partition coefficients between CaS and silicate melt. No rever-

Table 3. Experimental conditions, concentrations of REE in CaS and silicate, and REE CaS/silicate partition coefficients.

Run	CYA	CYC	CYB		CEI (rev.)	CEC	CEE	CEH (rev.)	CED ^a
T (°C)	1202	1306	1308		1202	1273	1295	1304	1308
Time(h)	18	21	20		18	16	16	22	26
wt%				Oldhamite					
Ca	51.80	48.67	47.46	Ca	49.31	50.31	49.91	48.53	52.97
Mg	1.76	4.4	4.83	Mg	0.11	2.27	2.40	3.62	0.46
Fe	0.30	0.92	1.04	Fe	0.37	0.69	0.88	0.92	1.88
Na	0.12	0.11	0.09	Na	0.10	0.08	0.09	0.19	0.09
S	42.36	43.52	43.17	S	43.64	41.23	41.70	44.75	44.58
La	0.39	0.15	0.21	Sm	0.75	0.53	0.55	0.79	0.42
Yb	0.43	0.46	0.52	Eu	0.83	0.68	0.78	1.04	0.49
Lu	0.39	0.42	0.45	Gd	0.71	0.48	0.51	0.99	0.41
sum ^b	97.55	98.65	97.77	sum ^b	95.82	96.27	96.82	100.8	101.3
wt%				Silicate matrix					
SiO ₂	52.04	45.39	47.23	SiO ₂	54.27	50.84	52.96	47.32	57.78
Al ₂ O ₃	11.03	21.43	19.71	Al ₂ O ₃	11.58	10.29	10.30	16.83	6.48
CaO	15.75	15.30	14.68	CaO	16.20	20.38	18.91	14.76	23.11
MgO	8.12	15.16	15.68	MgO	8.69	11.72	10.88	17.42	6.51
TiO ₂	0.79	0.31	0.32	TiO ₂	0.96	0.99	1.24	0.28	1.39
Na ₂ O	0.17	0.19	0.20	Na ₂ O	0.10	0.08	0.13	0.17	0.23
K ₂ O	0.18	0.16	0.18	K ₂ O	0.145	0.08	0.14	0.15	0.28
FeO	0.06	0.14	0.13	FeO	<0.03	0.18	0.14	0.10	0.59
La ₂ O ₃	3.68	0.44	0.46	Sm ₂ O ₃	2.15	1.83	1.88	0.75	0.89
Yb ₂ O ₃	3.39	0.37	0.36	Eu ₂ O ₃	0.52	0.68	0.80	0.56	0.50
Lu ₂ O ₃	3.35	0.40	0.42	Gd ₂ O ₃	2.29	1.69	1.76	0.92	0.98
sum	98.56	99.29	99.37	sum	96.91	98.76	99.14	99.26	98.74
D(La)	0.12	0.41	0.52	D(Sm)	0.45	0.34	0.34	1.23	0.55
D(Yb)	0.14	1.44	1.57	D(Eu)	1.87	1.13	1.14	2.16	1.14
D(Lu)	0.13	1.04	1.20	D(Gd)	0.36	0.33	0.34	1.24	0.48

Election Microprobe Analyses (LODDERS *et al.*, 1990). a: also contains coexisting FeS.

b: low totals due to surface oxidation. rev.: reversal experiment, REE initially contained in CaS.

sal experiments to test if equilibrium was achieved were published by JONES and BOYNTON (1983) or DICKINSON *et al.* (1990a, b, 1991). JONES and BOYNTON (1983) mention that their experimental charges probably are non-equilibrium assemblages, while DICKINSON *et al.* assume that equilibrium was reached between silicate liquid and immiscible sulfide liquid.

REE CaS/silicate partition coefficients are shown in Fig. 4. All groups investigating REE CaS/silicate partition coefficients found that partition coefficients increase from La to Lu (JONES and BOYNTON, 1983; DICKINSON *et al.*, 1990a, b, 1991; LODDERS *et al.*, 1990; LODDERS, 1991, 1995). However, absolute D values vary by about a factor of 20. These differences may be due to the different composition of the sulfide and silicate phases and may also be due to different oxygen and sulfur fugacities. The major difference in partitioning behavior was found for Eu. DICKINSON *et al.* (1990a, b, 1991) found that D_{Eu} for

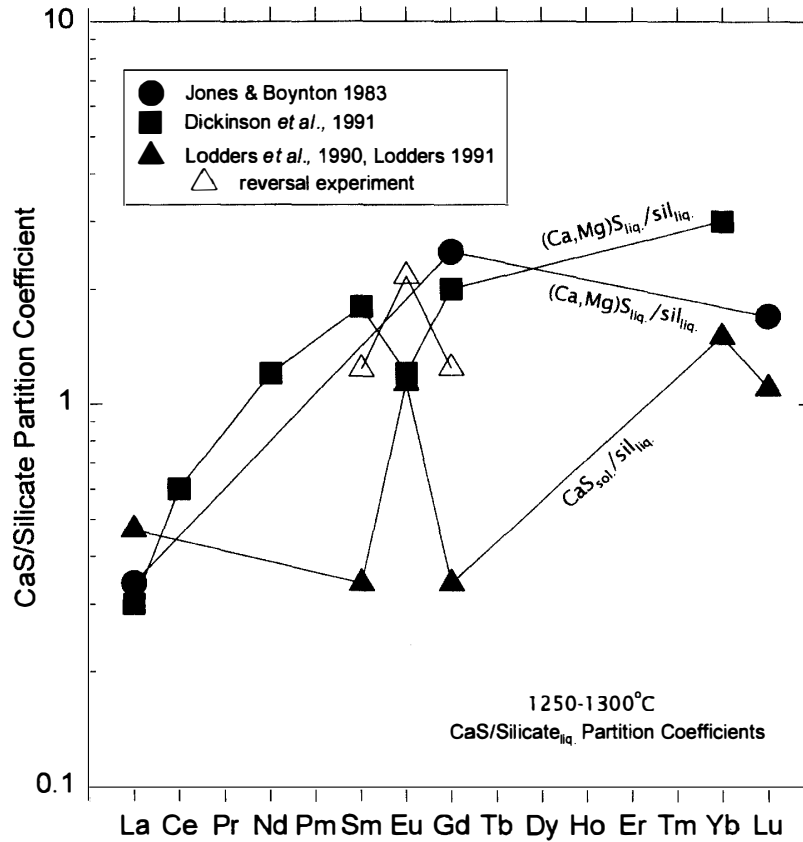


Fig. 4. *CaS/silicate partition coefficients for the REE at 1250–1300°C under reducing conditions. Experiments by JONES and BOYNTON and by DICKINSON et al. led to formation of a (Ca,Mg)S liquid coexisting with silicate melt while in experiments by LODDERS solid CaS coexisted with silicate melt. The Eu partition coefficient in the experiments by DICKINSON et al. is lower than that of neighboring Sm or Gd but the opposite is found in experiments by LODDERS. Reversal experiments (open symbols), where the REE had to move from CaS into silicate are 2–4 times higher than the partition coefficients obtained from experiments where REE initially were contained in the silicate, but also give $D_{Eu} > D_{Sm} \approx D_{Gd}$. This indicates that equilibrium was not reached, but the data from the reversal experiments give upper limits to the partition coefficients. No reversal experiments checking if equilibrium was obtained were published by JONES and BOYNTON or DICKINSON et al.*

(Ca, Mg)S/silicate was smaller than D values for neighboring Sm and Gd. This behavior of Eu is opposite to that found in the experiments by LODDERS *et al.* (1990) and LODDERS (1991), where $D_{Eu} > D_{Sm} \approx D_{Gd}$. The chemical behavior of Eu^{2+} is similar to that of Ca^{2+} and Eu may be preferentially incorporated into Ca-rich, Mg-poor sulfides over Mg-rich, Ca-poor sulfides. Experiments for REE FeS/silicate partitioning indicated that the partition coefficients depend on oxygen fugacity (see Fig. 3). DICKINSON *et al.* (1990b) also found that the partition coefficients for the CaS/silicate system seem to depend on oxygen fugacity. More experimental work is needed to investigate the REE partitioning as a

function of oxygen fugacity as well as of sulfide and silicate composition. However, all experimental results indicate that REE sulfide/silicate partition coefficients under reducing conditions are relatively small ($D < 10$).

4. REE Distribution in Aubrites: Where did Oldhamite Originate?

To test if aubritic minerals, including oldhamite, are equilibrated with respect to REE and thus formed by crystallization processes, the REE concentration ratios in the mineral phases are compared to the respective partition coefficient ratios. A comparison of observed and predicted ratios is given in Fig. 5. The observed ratios were calculated

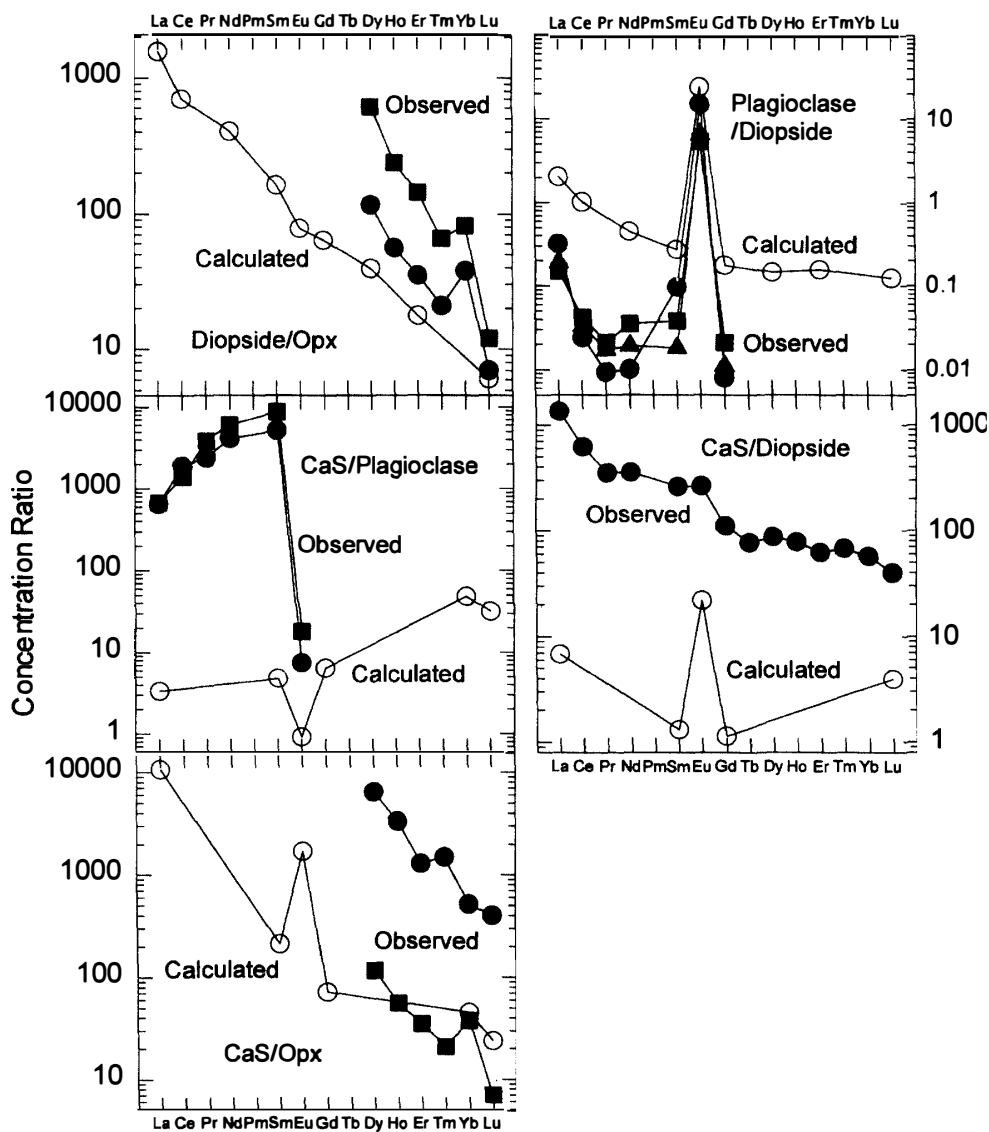


Fig. 5. Observed REE concentration ratios in mineral pairs in aubrites and calculated ratios from partition coefficients. Observed and calculated ratios do not match, indicating that no co-crystallization took place on the APB. For data sources and more discussion see text.

using the REE measured concentrations in aubritic mineral phases (data from FLOSS and CROZAZ, 1993; WHEELLOCK *et al.*, 1994). Although oldhamite in aubrites displays a variety of patterns, the oldhamite pattern displaying a negative Eu anomaly was selected because this was observed in an oldhamite dominated clast in the Norton County aubrite (WHEELLOCK *et al.*, 1994). In this clast, oldhamite is apparently in igneous contact to other aubritic minerals so that it provides a good test if oldhamite and other aubritic minerals are in equilibrium with respect to REE.

The predicted REE ratios were calculated from the respective mineral/silicate liquid partitioning data (KENNEDY *et al.*, 1993; GRUTZECK *et al.*, 1974; DRAKE and WEILL, 1975) and CaS/silicate partition coefficients from LODDERS *et al.* (1990) and LODDERS (1991, 1995). Although the CaS/silicate partition coefficients may not represent equilibrium assemblages, the results of the reversal experiments give upper limits to the CaS/silicate partition coefficients. In addition, all studies show REE CaS/silicate partition coefficients of the same order of magnitude.

The observed and calculated ratios shown in Fig. 5 disagree indicating that the REE bearing minerals are not equilibrated. Thus, co-crystallization from a parent melt can be excluded, as also discussed by GRAHAM and HENDERSON (1985). The other possible crystallization process is fractional crystallization and subsequent removal of crystallized phases from the melt (OKADA *et al.*, 1988). GRAHAM and HENDERSON (1985) concluded that a late crystallization of plagioclase in a fractional crystallization sequence is consistent with the REE distributions in the Mayo Belwa aubrite. However, the high REE concentrations in oldhamite are not explained by any of these crystallization processes.

In any crystallization sequence, the refractory (mp. 2450°C) oldhamite is expected to be among the first phases to crystallize, although it is not clear how CaS-rich oldhamite can crystallize from a Mg-rich melt. It has yet to be shown experimentally that Mg-poor oldhamite can form from an aubritic melt containing less than one percent Ca. The experiments by JONES and BOYNTON (1983) and DICKINSON *et al.* (1990a, b, 1991), where oldhamite was forced to form from a silicate melt by reaction with iron sulfide all yielded Mg-rich oldhamite.

Partition coefficients of about 100–200 are necessary to extract all REE from a melt into the small amount (~1 wt%) of oldhamite found in aubrites. However, the measured partition coefficients are about ≤ 10 . If we assume that enstatite and olivine crystallize and CaS continues to equilibrate with the remaining melt, REE partitioning into CaS is more efficient. However, even in this case, neither the observed REE abundances in CaS nor the abundance patterns can be explained by partitioning. One could argue that the experimentally determined partition coefficients are too small because equilibrium was not reached between the CaS and silicate melt. However, the reversal experiments gave D values within a factor of 2–4 of the D values obtained from experiments with REE-doped silicate. Furthermore, the work by JONES and BOYNTON (1983) and DICKINSON *et al.* (1990a, b, 1991) also gave partition coefficients $\ll 100$. Thus, models postulating the efficient incorporation of REE into oldhamite from a silicate liquid are not supported by the available experimental data and appear extremely unlikely.

Another problem with crystallization processes is that they cannot explain the different REE patterns in oldhamite or diopside. At least 10 different types of patterns have been observed for oldhamite and at least three different patterns have been reported for

diopside (FLOSS and CROZAZ, 1993). LODDERS and PALME (1990) proposed that oldhamite in aubrites is a relict phase from the solar nebula. The diversity of REE patterns in aubritic oldhamite was later seen as an indicator of relict origin of oldhamite by other groups (FLOSS *et al.*, 1990; KURAT *et al.*, 1992; FLOSS and CROZAZ, 1993). The REE abundances and REE patterns in oldhamite from aubrites are similar to those in enstatite chondrites, which can be explained by REE condensation into oldhamite from a highly reduced nebular gas (LARIMER *et al.*, 1984; LODDERS and FEGLEY, 1993). Other indicators of the condensation origin of aubritic oldhamite are the mass independent sulfur isotopic anomalies observed by THIEMENS *et al.* (1994) because crystallization processes would isotopically equilibrate the anomalous sulfur in oldhamite with all the other isotopically 'normal' sulfur in the whole rock.

WHEELOCK *et al.* (1994) proposed that oldhamite in a oldhamite-dominated clast in Norton County crystallized from a melt and that the high REE abundances in the CaS resulted from equilibration with the aubritic melt. This suggestion is favored by the large (up to 2 cm) oldhamite crystal size, which is not easily explained by condensation processes. WHEELOCK *et al.* postulated the presence of a liquid CaS phase because forsterite grains are surrounded by oldhamite, and because rounded sulfide blebs occur in oldhamite. However, temperatures of about 2450°C are required for *liquid* CaS and even if melting point depression of CaS due to dissolved MnS, FeS, or MgS is considered temperatures above 2000°C are necessary (LODDERS *et al.*, 1993). It is doubtful if these temperatures were reached on the APB. High temperatures would also lead to substantial vaporization of *e.g.*, Na, K, Si, and Mg (FEGLEY and CAMERON, 1987). However, evaporative loss did not occur because aubrites still contain moderately volatile elements such as Na and K. Thus it is questionable that temperatures on the APB were high enough to melt oldhamite.

5. Preferred Differentiation Model of the Aubrite Parent Body and the Origin of Oldhamite

The aubrite parent body assembled from material compositionally similar to EH-chondrites (FOGEL *et al.*, 1988; LODDERS *et al.*, 1993). The REE are mainly located in oldhamite, which gained REE during condensation, as reflected by the diversity of its REE patterns and the high REE abundances. Impact melting led to heating and differentiation of the APB. Once temperatures reached ~1000°C, the Fe-FeS eutectic started segregating to form a core. Because oldhamite contains the REE and is not expected to be involved in major exchange reactions at these temperatures, the segregating FeS did not take up any significant amounts of REE. The low density of CaS (2.5 g/cm³) allowed flotation and accumulation of small CaS grains once the silicates started melting. An example of such an accumulation process may be the CaS-rich 1.6 cm long impact melt vein in the Jajh deh Kot Lalu (EL6)-enstatite chondrite (RUBIN *et al.*, 1995). Once larger concentrations of CaS were produced, sintering led to larger CaS aggregates. Formation of larger CaS aggregates by sintering was observed in CaS/silicate partitioning experiments (LODDERS *et al.*, 1990; LODDERS, 1991). However, not all oldhamite inherited from the enstatite chondrite-like precursor may have come in contact with melt and accumulated. Most oldhamite in aubrites is small and comparable in grain size with that in enstatite chondrites (30–200 μm ; LARIMER and GANAPATHY, 1987).

LODDERS *et al.* (1993) previously proposed that the different REE patterns in diopside can be understood if diopside was formed from oldhamite. FOGEL *et al.* (1988) suggested the reaction of oldhamite with enstatite to form diopside and forsterite. In that case, the diopside would inherit the REE inventory of the oldhamite. FLOSS and CROZAZ (1993) report REE patterns in diopside, which contains a weathered oldhamite inclusion. This diopside may be the reaction product from oldhamite with enstatite, because its REE pattern is unlike those found in other diopsides in aubrites. This diopside shows two patterns with light REE around 0.5 times chondritic and heavy REE around 1 times chondritic. In addition, one pattern displays a small negative Eu anomaly. The oldhamite inclusion has a relatively flat pattern about 100 times chondritic. If diopside formed solely by crystallization from a melt, only the REE pattern expected from diopside/melt partitioning should be observed, instead of the different types of patterns which are actually found.

Crystallization processes also cannot explain the observed plagioclase zoning with anorthite at the rim and albite in the center, which was described by WHELOCK *et al.* (1994) for an oldhamite dominated clast in the Norton County aubrite. Anorthite is more refractory than albite and crystallizes first. Thus a crystallization process predicts an anorthite rich interior and albitic rim instead. However, reaction of oldhamite with albite may explain the inverse zoning and also explain the negative Eu-anomaly in some aubritic oldhamites, because plagioclase favors the uptake of Eu over other REE.

It appears that the oldhamite from the precursor material of aubrites underwent different degrees of exchange reactions with its surrounding minerals but that these exchange reactions never reached equilibrium. These exchange process should also lead to zoning profiles in the REE distributions among minerals coexisting with CaS. Further REE measurements on such mineral assemblages may reveal that the REE underwent metamorphic redistribution. For example, preliminary ion probe investigations of enstatite in contact with oldhamite from the Pena Blanca Spring aubrite (FAHEY *et al.*, 1995) show a falling concentration gradient of the REE from the oldhamite-enstatite interface towards the enstatite interior. Clearly more analytical work on aubrites and experimental work for modeling the behavior of REE among aubritic minerals is needed to understand the differentiation history of the APB.

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