

A hibonite-corundum inclusion from Murchison: A first-generation condensate from the solar nebula

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Abstract-Through freeze-thaw disaggregation of the Murchison (CM) carbonaceous chondrite, we have recovered a $\sim 90 \times 75 \ \mu m$ refractory inclusion that consists of corundum and hibonite with minor perovskite. Corundum occurs as small (~10 μ m), rounded grains enclosed in hibonite laths (~10 μ m wide and 30–40 μ m long) throughout the inclusion. Perovskite predominantly occurs near the edge of the inclusion. The crystallization sequence inferred petrographically-corundum followed by hibonite followed by perovskite—is that predicted for the first phases to form by equilibrium condensation from a solar gas for $P^{\text{tot}} \leq 5 \times 10^{-3}$ atm. In addition, the texture of the inclusion, with angular voids between subhedral hibonite laths and plates, is also consistent with formation of the inclusion by condensation. Hibonite has heavy rare earth element (REE) abundances of $\sim 40 \times CI$ chondrites, light REE abundances ~20 × CI chondrites, and negative Eu anomalies. The chondritenormalized abundance patterns, especially one for a hibonite-perovskite spot, are quite similar to the patterns of calculated solid/gas partition coefficients for hibonite and perovskite at 10^{-3} atm and are not consistent with formation of the inclusion by closed-system fractional crystallization. In contrast with the features that are consistent with a condensation origin, there are problems with any model for the formation of this inclusion that includes a molten stage, relic grains, or volatilization. If thermodynamic models of equilibrium condensation are correct, then this inclusion formed at pressures $<5 \times 10^{-3}$ atm, possibly with enrichments ($<1000 \times$) in CI dust relative to gas at low pressures (below 10-4 atm). Both hibonite and corundum have $\delta^{17}O \approx \delta^{18}O \approx -50\%$, indicating formation from an ¹⁶O-rich source. The inclusion does not contain radiogenic ²⁶Mg and apparently did not contain live 26 Al when it formed. If the short-lived radionuclides were formed in a supernova and injected into the early solar nebula, models of this process suggest that ²⁶Al-free refractory inclusions such as this one formed within the first $\sim 6 \times 10^5$ years of nebular collapse.

INTRODUCTION

Equilibrium thermodynamic condensation calculations for a cooling solar gas at $P^{\text{tot}} < 10^{-2}$ atm (*e.g.*, Yoneda and Grossman, 1995) show that corundum should be the first major condensate. With continued cooling, corundum is predicted to react with the nebular vapor to form hibonite. Apparently, in many cases it did so, because hibonite-bearing inclusions are much more abundant than corundum-bearing ones in carbonaceous chondrites. Inclusions containing possible primary corundum and hibonite have been reported by Bar-Matthews *et al.* (1982), Fahey (1988) and by Krot *et al.* (2001). In their study of BB-5, the first corundum-bearing inclusion ever reported, Bar-Matthews *et al.* (1982) favored a condensation origin over an igneous one, but could not draw a firm conclusion because (a) they could not tell from the texture of the sample whether it crystallized from a melt or was a direct gas-solid condensate, and (b) adequate thermodynamic data for hibonite were not yet available. It was not known whether hibonite should condense from a gas of solar composition, and, if so, whether it should do so before or after corundum. The corundum in another inclusion, GR-1, most likely formed by the partial breakdown of hibonite (MacPherson *et al.*, 1984; Hinton *et al.*, 1988). Hibonite, being the most refractory major phase that is a carrier of rare earth elements (REE), and commonly containing isotopic anomalies (*e.g.*, Ireland *et al.*, 1991), can yield important insights into physico-chemical conditions in the early solar nebula.

The Murchison inclusion described herein appears to contain primary corundum enclosed in hibonite. With the

RESULTS

microprobe, we have conducted a detailed study of this object in order to gain insights into the early history of the solar system. Pristine, unmelted inclusions are especially important in light of the growing number of studies (*e.g.*, MacPherson and Davis, 1993; Beckett *et al.*, 2000; Grossman *et al.*, 2000) that provide strong evidence that many refractory inclusions, especially type B inclusions, have undergone complex thermal histories involving, for example, multiple melting events and probably evaporation. Such events can overprint the chemical and isotopic information recorded earliest in the inclusions, whereas unmelted condensate assemblages potentially provide direct information about possible pressures, temperatures, and isotopic compositions in the early solar nebula. Preliminary results of this study were reported by Simon *et al.* (2000a).

scanning electron microscope, electron microprobe and ion

ANALYTICAL METHODS

The sample, M98-8, was recovered by hand-picking from the high-density ($\rho > 3.2$) separate obtained from the products of freeze-thaw disaggregation of a bulk sample of Murchison by the method of MacPherson et al. (1980). Backscattered electron images and x-ray maps of a polished thin section of the sample were obtained with a JEOL JSM-5800LV scanning electron microscope equipped with an Oxford/Link ISIS-300 energy-dispersive x-ray microanalysis system. Quantitative wavelength-dispersive analyses were obtained with a Cameca SX-50 electron microprobe operated at 15 kV with a beam current of 25 nA. Data were reduced via the modified ZAF correction procedure PAP (Pouchou and Pichoir, 1984). Trace element and Mg isotopic analyses were obtained using the University of Chicago AEI IM-20 ion microprobe. The analytical techniques used are similar to those described in Simon et al. (1991), MacPherson and Davis (1993, 1994) and Russell et al. (2000). Trace element analyses were done using energy filtering. A variety of silicate standards were used to determine calcium-normalized yields. NIST 611 glass was analyzed at the beginning and end of the run to correct for variations in ion yield that are a function of mass (MacPherson and Davis, 1994). For Mg isotopic analyses, Madagascar hibonite was used as a standard for determining instrumental mass fractionation.

After cleaning and recoating of the sample, oxygen isotopic abundances were determined with the UCLA CAMECA ims 1270 ion microprobe utilizing techniques similar to those described in Simon *et al.* (2000b). A Cs primary ion beam was used to sputter shallow, elliptically-shaped craters of ~12×18 μ m. Analyses were performed without energy filtering at high mass resolving power using an electron flood gun for achieving charge compensation. Sample topography and previous ion probe analyses limited the surface area available for oxygen isotope measurements and contributed to slightly larger errors of 2 to 3‰ (1 σ) for δ ¹⁸O and δ ¹⁷O in each of three spots analyzed.

Petrography

The sample appeared pale green prior to sectioning but is colorless in thin section. A backscattered electron image of M98-8 (Fig. 1) shows that it is rounded, and $\sim 90 \times 75 \,\mu$ m. It is dominated by subhedral hibonite laths that are mostly $\sim 10 \,\mu m$ wide and $30-40\,\mu\text{m}$ long. Most hibonite laths enclose rounded, anhedral grains of corundum that are 5 μ m across. The largest corundum grain is ~15 μ m in size. Small grains of perovskite, a few microns across, are also present, mostly occurring at the edges of hibonite grains and at the edge of the inclusion. Small grains of Mg-Fe phyllosilicate, a secondary phase that is common in Murchison, are found predominantly at the edge of the inclusion, but also in some of the interior cavities. Elemental x-ray maps (Fig. 2) help illustrate the distribution of phases. The phyllosilicate is the bright phase in the Si map, and perovskite is bright in the Ca and Ti maps. Corundum is bright in the Al map and black in the Ca map. From counting 5870 points (excluding the voids) on the backscattered electron image, we obtained a mode of 89.1 vol% hibonite, 9.3% corundum, and 1.6% perovskite.

There is void space between many of the hibonite grains, giving the object a somewhat fluffy texture. Many of these voids, their shapes controlled by hibonite crystals, are triangular- or trapezoidal-shaped gaps between laths, like those shown in Fig. 3. In some of the voids, hibonite crystal faces can be seen jutting into open space, and some contain unpolished plates below the surface of the section (Fig. 3b). The textures of two corundum-bearing inclusions from Adelaide can also be described as fluffy. They were found in situ (Krot et al., 2001), and meteorite matrix or rim material occurs between many of the hibonite crystals. The overall texture of M98-8 is different from those of the two other hibonitecorundum inclusions that have been found in Murchison, BB-5 (Bar-Matthews et al., 1982) and GR-1 (MacPherson et al., 1984). Sample BB-5 is more compact than M98-8 and its corundum is subhedral and concentrated in the core of the inclusion, rather than dispersed throughout the inclusion as in M98-8. Sample GR-1 has a euhedral hibonite crystal at its core. This grain is enclosed in corundum, and hibonite is also present at the edge of the inclusion. Murray sample F5 (Fahey, 1988) is similar to the Adelaide inclusions in that it consists of hibonite laths that enclose small corundum grains and appears to be very loosely consolidated, with meteorite matrix between hibonite grains.

Mineral Chemistry

Corundum compositions are nearly pure Al_2O_3 , as shown by the representative analyses given in Table 1. Small amounts of TiO₂ are also present, as is the case for BB-5 (Bar-Matthews *et al.*, 1982) and GR-1 (MacPherson *et al.*, 1984). Corundum



FIG. 1. Backscattered electron image of M98-8. Corundum is the dark gray phase seen at the upper left and as isolated, rounded grains enclosed in hibonite (light gray). Void spaces and epoxy are black. cor = corundum; hib = hibonite; pv = perovskite.

Table 1.	Electron microprobe analyses of
corundum	in M98-8.

	1	2		
Al ₂ O ₃	99.85	99.61		
SiO ₂	0.02	BDL		
CaO	0.09	0.05		
TiO ₂	0.17	0.25		
FeO	BDL	0.04		
Total	100.13	99.95		
Cations per 3 oxygen anions				
Al	1.995	1.995		
Si	0	0		
Ca	0.002	0.001		
Ti	0.002	0.003		
Fe	0	0.001		
Total cations	1.999	2.000		

 $\begin{array}{l} BDL = below \mbox{ detection limit (in wt%) of } 0.012 \mbox{ for } SiO_2; \mbox{ or } 0.036 \mbox{ for } FeO. \mbox{ Also below detection:} \\ MgO \ (<\!0.015 \ wt\%); \ Sc_2O_3 \ (<\!0.032); \mbox{ and } V_2O_3 \ (<\!0.027). \end{array}$

in F5 contains <0.1 wt% TiO₂ (Fahey, 1988). In the M98-8 corundum, as in the other occurrences, abundances of MgO, Sc_2O_3 , and V_2O_3 are all below the detection limits of the electron microprobe.

Representative compositions of hibonite from M98-8 are given in Table 2. The crystals are not zoned, and all contain <2 wt% TiO₂. The overall range in TiO₂ contents is from 1.2 to 1.9 wt%, and most grains have between 1.5 and 1.9 wt%. Abundances of Sc₂O₃, V₂O₃, FeO and, in most cases, SiO₂ are below the detection limits of the electron probe. Hibonite in the other corundum-bearing inclusions also has TiO2 contents ≤ 2 wt%. The large, single crystals of hibonite found in Murchison also tend to have low TiO2 contents, while the hibonite from spinel-hibonite spherules typically has 3-7 wt% TiO₂ (Ireland, 1988). Trivalent Ti can substitute for Al in hibonite, and tetravalent Ti and Si cations can enter hibonite along with Mg via a coupled substitution for two Al³⁺ cations. If Ti³⁺ proportions are negligible, a plot of Mg vs. Ti + Si cation abundances in meteoritic hibonite typically yields a trend with a slope of 1. Such a line, shown for reference in Fig. 4, passes through the data, with most points on or close to the line, suggesting that Ti³⁺ proportions are low. If Ti³⁺ contents were significant, the data would plot on the high-Ti



FIG. 2. Backscattered electron image (upper left) and elemental x-ray maps of M98-8. Note the distribution of corundum (bright in Al map) throughout the inclusion, and the tendency of perovskite (bright in Ca and Ti maps) and phyllosilicate (bright in Si map) to occur at the edge of the inclusion.



FIG. 3. Secondary electron images of M98-8 showing void space bounded by crystal faces (*e.g.*, upper right in both (a) and (b)) and unpolished plates below the surface of the section (arrow near center of (b)).



Ti + Si cations per 19 Ox

FIG. 4. Electron microprobe analyses of hibonite in M98-8. The hibonite has low TiO_2 contents and the data scatter about the 1:1 correlation line for Mg vs. Ti + Si cations, which is shown for reference.

TABLE 2. Electron microprobe analyses of hibonite in M98-8.

	1	2	3
MgO	0.81	0.98	0.90
Al_2O_3	88.96	88.86	88.61
SiO ₂	BDL	0.08	BDL
CaO	8.59	8.59	8.42
TiO ₂	1.64	1.88	1.92
Total	100.00	100.39	99.85
	Cations pe	er 19 oxygen anions	
Mg	0.134	0.162	0.150
Al	11.706	11.651	11.676
Si	0	0.009	0
Ca	1.028	1.024	1.009
Ti	0.138	0.158	0.161
Total cations	13.006	13.004	12.996

BDL = below detection limit of 0.012 wt%. Also below detection: Sc_2O_3 (<0.042 wt%); V_2O_3 (<0.033); and FeO (<0.039).

side of the slope-1 line and further from it with increasing Ti content.

Trace Element Abundances

Ion microprobe analyses are given in Table 3, and chondritenormalized REE abundance patterns are illustrated in Fig. 5. All of the spots are hibonite-dominated, though one spot sampled perovskite (probably ~ 4 wt%, based on the TiO₂ content) and Mg-, Fe-silicate in addition to hibonite. The other two spots have rather similar trace element concentrations to each other, so they were averaged for Fig. 5. The patterns have some features in common, and some differences. Each of the analyses has (chondrite-normalized) heavy REE (HREE) > light REE (LREE), as found in ultrarefractory inclusions, although the HREE enrichment relative to the LREE in M98-8 is not as pronounced as it is in most ultrarefractory inclusions. The pattern for the perovskite-poor analysis has flat LREE abundances as does the perovskite-rich analysis, except that abundances decrease from Nd through Eu in the latter. The chondrite-normalized HREE patterns are different from each other and are unusual in that Tm is enriched relative to Er in



FIG. 5. Chondrite-normalized rare earth element and refractory trace element abundances in M98-8, measured by ion microprobe. The enrichment of heavy REE relative to light REE is characteristic of ultrarefractory inclusions.

one analysis and enriched relative to Lu in the other, whereas in most ultrarefractory inclusions Tm is significantly depleted relative to both Er and Lu. The hibonite-corundum pattern, in which Tb, Dy, Y and Ho have the greatest enrichment factors with a "rolloff" through the heavier REE resembles the hibonite + perovskite pattern reported for BB-5 (Hinton et al., 1988). In contrast, the Murray inclusion F5 has a group II REE pattern (Fahey, 1988). Among other trace elements, the hibonite in M98-8 contains ~400 ppm Sc and 120-350 ppm Zr, and these values are within the range of those reported for Murchison hibonite by Ireland et al. (1988). The CI-normalized refractory trace element abundance pattern for the perovskite-poor analysis is similar to that for the perovskite-rich one, except that the former is depleted in Mo and Nb relative to the latter (Fig. 5). This is not surprising, as perovskite is typically Nb-rich (Ireland et al., 1988).

Isotopic Abundances

Measurements of the intrinsic mass-fractionation of magnesium, F_{Mg} , are all within 2σ error of -5%, and three of

the five analyses are within error of 0‰ (Table 4). Neither the hibonite nor the corundum incorporated measurable live ²⁶Al when they formed. On a plot of δ^{26} Mg vs. ²⁷Al/²⁴Mg (Fig. 6), the best-fit line through the data has a negative slope, but all the measurements of δ^{26} Mg are within error of 0. Many calcium-aluminum-rich inclusions (CAIs) consist of phases with excess ²⁶Mg abundances that are consistent with initial ²⁶Al/²⁷Al ratios of ~5 × 10⁻⁵ (*e.g.*, MacPherson *et al.*, 1995). In contrast, the upper limit for the initial ²⁶Al/²⁷Al ratio of the present sample, based on the 2 σ upper limit on the slope of the line through the data points, is ~1.6 × 10⁻⁶. No evidence of live ²⁶Al was found in BB-5 (Bar-Matthews *et al.*, 1982), while Fahey (1988) reported an initial ²⁶Al/²⁷Al ratio of (4.1 ± 0.2) × 10⁻⁵ for F5.

Oxygen isotopic compositions in M98-8 were determined in two spots of pure hibonite, while spot three sampled a mixture of hibonite and corundum (Table 5). We do not know precisely the corundum/hibonite ratio of the analytical volume, but inspection of the spot three pit after analysis indicates that it apparently contained at least 50% hibonite. As the composition of the corundum + hibonite spot is within error of the hibonite analyses, the isotopic composition of the corundum is probably

TABLE 3. Major and trace element abundances in three analysisspots on Murchison M98-8, determined by ion microprobe.*

	#6	#7	#8
Phase [†]	hib-cor	hib-pv-sil	hib-cor
MgO	0.701	1.39	0.766
Al ₂ O ₃	87.2	72.6	88.0
SiO ₂	0.243	10.1	0.527
CaO	9.58	9.02	8.28
TiO2	1.97	4.32	1.81
FeO	0.213	1.27	0.440
т:	0.254 ± 0.054	0.95 ± 0.12	0.200 ± 0.051
LI	0.234 ± 0.034	0.83 ± 0.12 0.78 ± 0.11	0.200 ± 0.031
De	0.848 ± 0.093	0.78 ± 0.11	0.87 ± 0.10
В	0.72 ± 0.18	5.10 ± 0.58	1.55 ± 0.29
Na	45.7	5930	104
P	5.4 ± 3.9	<36	11.7 ± 6.0
ĸ	11.6 ± 1.3	341	24.9 ± 2.3
Sc	393	424	461
V	70.1 ± 5.2	68.8 ± 7.6	63.3 ± 5.6
Cr	5.7 ± 2.0	149 ± 9	44.5 ± 4.6
Mn	9.0 ± 1.6	27 ± 14	15.4 ± 2.5
Co	4.6 ± 2.2	40.6 ± 7.3	7.0 ± 2.9
Ni	97 ± 16	896 ± 55	185 ± 25
Rb	0.34 ± 0.25	2.19 ± 0.75	0.35 ± 0.26
Sr	38.1	67.0	46.0
Y	71.6	151	55.6
Zr	138	351	123
Nb	0.088 ± 0.083	10.4 ± 1.1	0.32 ± 0.18
Mo	2.4 ± 1.1	39.8 ± 5.8	8.0 ± 2.3
Ba	8.35 ± 0.64	9.60 ± 0.86	8.09 ± 0.70
La	3.44 ± 0.25	9.16 ± 0.50	4.66 ± 0.32
Ce	6.19 ± 0.54	25.1 ± 1.3	10.6 ± 0.8
Pr	1.21 ± 0.29	4.22 ± 0.66	1.06 ± 0.30
Nd	6.8 ± 1.6	20.1 ± 3.3	4.3 ± 1.4
Sm	1.89 ± 0.50	3.51 ± 0.82	1.70 ± 0.51
Eu	0.27 ± 0.13	0.56 ± 0.22	0.40 ± 0.15
Gd	5.7 ± 1.2	5.3 ± 1.8	6.4 ± 1.3
Tb	1.54 ± 0.22	1.11 ± 0.28	1.66 ± 0.25
Dy	11.1 ± 1.4	15.7 ± 2.1	6.2 ± 1.2
Ho	2.31 ± 0.31	3.66 ± 0.49	1.84 ± 0.31
Er	5.24 ± 0.83	19.8 ± 2.0	2.91 ± 0.68
Tm	0.62 ± 0.11	4.71 ± 0.36	0.338 ± 0.090
Yb	0.48 ± 0.43	1.58 ± 0.70	< 0.76
Lu	< 0.20	8.98 ± 0.51	< 0.22
Th	0.56 ± 0.15	0.77 ± 0.21	0.80 ± 0.19
U	0.115 ± 0.066	< 0.11	< 0.089

*Oxide concentrations are given in wt% and elemental concentrations in ppm. Uncertainties are $\pm 1\sigma$, based on counting statistics, and are only given where they exceed 5% of the amount present; upper limits are $< 2\sigma$.

[†]Abbreviations: hib = hibonite; cor = corundum; sil = Mg, Fe-rich silicate.

TABLE 4. ²⁷Al/²⁴Mg ratios and Mg isotopic compositions of M98-8.

	27Al/24Mg	$F_{\rm Mg}$ (‰/amu)	$\delta^{26} { m Mg}$ (‰)
hib	113.6 ± 8.1	-10.2 ± 4.7	2.6 ± 6.8
hib	168.6 ± 3.3	-6.5 ± 3.3	-0.4 ± 5.0
hib	215 ± 27	-3.0 ± 3.8	-1.6 ± 4.9
hib	80.9 ± 0.6	-2.8 ± 3.9	-0.9 ± 5.2
cor	550 ± 73	-4.7 ± 7.1	-6.1 ± 11.4

Uncertainties are $\pm 2\sigma$. Abbreviations: hib = hibonite, cor = corundum.

similar to that of the hibonite. Both phases have ¹⁶O-rich compositions, similar to some that have previously been reported for CM hibonite (*e.g.*, Fahey *et al.*, 1987; Ireland *et al.*, 1992). These results indicate that M98-8 formed in an ¹⁶O-rich environment and, unlike many melilite-rich inclusions (Clayton *et al.*, 1977), did not partially re-equilibrate with an ¹⁶O-poor vapor following crystallization.

DISCUSSION

Formation of the Inclusion

We now consider ways in which this inclusion could have formed: gas-to-solid condensation, or crystallization from (a) a stable, condensate liquid; (b) a metastable liquid; or (c) a liquid formed by melting and/or evaporation of hibonite.

In thermodynamic models of equilibrium condensation from a solar gas for total pressures <10-2 atm, corundum is the first major condensate. At $P^{\text{tot}} = 10^{-3}$ atm, for example, corundum condenses at 1770 K and, with decreasing temperature, is predicted to react with the nebular gas to form hibonite at 1743 K, followed by condensation of perovskite at 1688 K (Yoneda and Grossman, 1995). At higher pressures ($\geq 10^{-2}$ atm), either hibonite, grossite, or liquid is the first condensate. With corundum grains enclosed in hibonite, and perovskite mostly occurring at the edge of the inclusion, the crystallization sequence consistent with the texture of M98-8 is the same as the equilibrium condensation sequence for a solar gas at low pressures. The rounded shapes of the corundum grains are consistent with the predicted corundum-hibonite reaction relationship, and the angular voids between hibonite grains also might be expected for an aggregate of condensate grains. For crystallization from a liquid, on the other hand, voids tend to be round and crystals tightly intergrown, with the shapes of late crystals conforming to the shapes of early crystals. These features are typically found in the relatively common hibonitespinel spherules (e.g., MacPherson et al., 1983, 1984).

Simon *et al.* (1996) calculated solid/gas distribution coefficients, summarized in Fig. 7a, for REE and Y condensation into hibonite and perovskite from the volatilities of these elements and activity coefficients for their solid solution into those phases. The chondrite-normalized REE abundance



FIG. 6. δ^{26} Mg, corrected for mass-fractionation, *vs.* ²⁷Al/²⁴Mg for hibonite and corundum in M98-8. There is no evidence for excess ²⁶Mg. A best-fit line through the data points has a *y*-intercept and a negative slope that are within error of 0. Lines for initial ²⁶Al/²⁷Al ratios of 0 and 5 × 10⁻⁵ are shown for reference. All uncertainties are ±2 σ .

	δ ¹⁸ O (‰)	2σ mean (‰)	δ ¹⁷ O (‰)	2σ mean (‰)	Δ ¹⁷ O (‰)	2σ mean (‰)	16O-excess (‰)	2σ mean (‰)
Spot 1	-55.8	4.3	-53.6	3.9	-24.6	3.1	51.2	6.5
Spot 2	-49.1	4.5	-47.4	4.2	-21.9	3.6	45.6	7.5
Spot 3	-54.5	6.1	-52.7	5.0	-24.3	4.9	50.6	10.3

TABLE 5. Oxygen isotopic compositions of phases in M98-8.

Spots 1 and 2 are hibonite. Spot 3 is corundum + hibonite. The mixing proportions are not well determined. The deviation of the oxygen isotopic composition from the terrestrial fractionation line may be expressed by $\Delta^{17}O$, where: $\Delta^{17}O = \delta^{17}O - 0.52 \times \delta^{18}O$. Alternatively, the distance along the CAI-mixing line (Clayton *et al.*, 1977) can be expressed by the ¹⁶O-excess (Clayton and Mayeda, 1983) given by the expression: $(0.52 \times \delta^{18}O - \delta^{17}O)/(1 - 0.52)$.

patterns of these phases in M98-8 are consistent with a condensation origin in that they qualitatively resemble these calculated solid/gas distribution coefficient patterns fairly closely while showing little similarity to the patterns of hibonite/ liquid or perovskite/liquid distribution coefficients measured by Kennedy *et al.* (1994). As shown in Fig. 7a, for both hibonite and perovskite, the solid/gas distribution coefficients decrease

from Lu through Sm, are fairly flat for the LREE, and are very low for Eu and Yb, in a pattern quite like that of the chondritenormalized REE abundances in the perovskite-rich analysis spot (Fig. 5). The only mismatch is for Tm, which, assuming a chondritic source, should be depleted relative to Er and have a chondrite-normalized abundance similar to those of the LREE but, in the sample, is enriched relative to Er and the LREE. It



FIG. 7. (a) Solid/gas distribution coefficients, relative to that for Lu, calculated for condensation of REE from a solar gas into perovskite and hibonite at the conditions shown. After Simon *et al.* (1996). Note resemblance of these patterns to the observed hibonite-perovskite pattern (Fig. 5). (b) Solid/liquid distribution coefficients for perovskite and hibonite from the experiments of Kennedy *et al.* (1994).

is difficult to account for the observed Tm abundance for this spot. Oxidizing conditions can cause Tm enrichment in the solid, but such conditions would lead to negative Ce anomalies (Davis *et al.*, 1982), and these are not observed. In contrast with the solid/gas distribution coefficients and with the analyses of M98-8, the crystal/liquid distribution coefficients decrease from La through Lu (Fig. 7b).

From the bulk composition of M98-8, we know that if the inclusion had crystallized from a melt, perovskite could only crystallize late, after most or all of the hibonite. The results of Kennedy et al. (1994) show, however, that the light REE are compatible in hibonite, so perovskite that crystallized after hibonite should have relatively low LREE contents. We estimate that the perovskite-bearing analysis (#7, Table 3) sampled ~4 wt% perovskite. Assuming that the REE contents of the silicate are negligible and that the La content of the hibonite is $\sim 18 \times CI$ (Fig. 5), we calculate that the La content of the perovskite is $>550 \times CI$. From the modal proportions of corundum, hibonite and perovskite in the inclusion, this leads to a bulk La content of $\sim 25 \times CI$. Thus, the perovskite has a La content much greater than the bulk, which is inconsistent with prior crystallization of ~90% of a phase with a crystal/liquid distribution coefficient for La of ~6 (Kennedy et al., 1994). The perovskite and the hibonite could not have crystallized from a common parental liquid.

There are additional problems with any model that includes a molten stage. From known phase relations (Berman, 1983) and the bulk Al₂O₃ content of M98-8 (over 88 wt%), we estimate that the inclusion would be completely molten only at temperatures >2140 K, but no condensed phases are stable in a solar gas at such temperatures at any $P^{\text{tot}} < 10$ atm (Yoneda and Grossman, 1995). For such an aluminous liquid to be stable at more reasonable total pressures, high dust/gas ratios and temperatures are required (e.g., ~1000 × solar and 2400 K, respectively, for $P^{\text{tot}} = 10^{-3}$ atm; Yoneda and Grossman, 1995). Even if such conditions could be attained, Ca would continue to condense into the liquid, decreasing its Al₂O₃ content, during cooling so that, at the liquidus temperature, the liquid would be too CaO-rich to crystallize corundum, and would crystallize hibonite instead. Thus, in order to crystallize corundum from a stable liquid in the solar nebula, extraordinary P-T conditions would be required to form an extremely Al₂O₃-rich, CaO-poor liquid (note that these temperatures are much higher than those required for direct condensation of corundum), and the very early condensate liquid would have to be isolated from the nebular vapor and allowed to cool without further reaction with Furthermore, although this scenario would yield it. crystallization of corundum followed by hibonite, it probably would not produce the observed texture. Molten droplets typically cool by radiating heat from their surfaces, a process which, in this case, would probably form an outer zone of corundum enclosing a core of hibonite. Instead, in M98-8, we find small corundum grains enclosed within individual crystals of hibonite. Formation of M98-8 by crystallization of a stable molten droplet thus seems highly unlikely.

Metastable liquids could have existed in the nebula. Experiments (Nelson et al., 1972; Keil et al., 1973) have shown, however, that during rapid cooling and solidification of supercooled alumina droplets, nucleation and crystal growth are rapid, yielding spherulitic, cryptocrystalline textures quite unlike the texture of M98-8, with its massive, well-formed hibonite crystals. Even slow cooling of such droplets would be expected to yield quench textures. Calculations by Nelson et al. (1972) suggest that, in a completely molten droplet, the absence of crystal nucleii could permit significant degrees of undercooling even at much slower cooling rates than those used in their experiments, leading to rapid crystallization and therefore textures similar to those of quenched spherules. Slow cooling would also allow extensive evaporation from the melt, which leads to problems as described below. The features of M98-8 are not consistent with crystallization from a completely molten, stable or metastable condensate liquid.

Another way corundum can form is through the breakdown of hibonite. Hibonite melts incongruently to corundum + liquid. If a hibonite-rich inclusion were to be heated, melted, and partially evaporated, a corundum-bearing residue could result. This may have occurred during the formation of GR-1, a hibonite-corundum inclusion from Murchison first described by MacPherson et al. (1984). That inclusion has a basically concentric structure, with hibonite at the center enclosed in corundum, and hibonite also at the outer edge of the inclusion. MacPherson et al. (1984) noted that the core and rim hibonite had different Ti, Mg and Sc contents, and Hinton et al. (1988) found that they have contrasting trace element and isotopic compositions as well. As noted by these workers, the texture and the chemical data strongly suggest that this inclusion contains two generations of hibonite. What appears to have happened is that a hibonite-rich object was partially melted, leaving some residual hibonite; corundum formed, enclosing the relic hibonite in the interior, and all of the Ca evaporated from the liquid. The corundum later reacted with a Ca-bearing vapor to form the outer, REE-poor hibonite (Hinton et al., 1988). Unlike that of M98-8, the texture of GR-1 suggests a hibonitecorundum-hibonite crystallization sequence, reflecting a complex thermal history.

Evaporation of hibonite was investigated experimentally by Floss *et al.* (1998), who rapidly heated, melted, and cooled hibonite samples in a vacuum furnace. They found that in all of their experiments, even one with only 6% mass loss, hibonite broke down completely, leading to run products consisting of corundum + glass \pm thorianite \pm a REE-rich phase. Unlike the GR-1 case, no relic hibonite was preserved, and no occurrences of corundum + hibonite were found. The occurrence of glass in these run products was probably due to the rapid cooling experienced by the liquid. If the melt had cooled slowly, avoiding formation of glass, it would have remained in the liquid state longer and presumably undergone fairly extensive evaporation of Mg and Ca. This would have resulted in strong isotopic mass fractionation, which is not observed, at least for Mg, in M98-8. In addition, evaporation of all oxides more volatile than Al_2O_3 would have lead to absence of hibonite.

In summary, there are serious problems with any model for the formation of M98-8 that includes a molten stage, relic phases or evaporation. Gas-to-solid condensation is the most straightforward way to account for the textural and chemical features of this sample.

Constraints on the Formation Conditions of M98-8

Given that M98-8 records a corundum-hibonite-perovskite (CHP) condensation sequence, thermodynamic calculations can be used to estimate under what combinations of temperature, pressure, and dust/gas ratio the sample could have formed. Increasing the dust/gas ratio increases the proportion of condensable elements in the system, and its effect is similar to that of increasing pressure. The model of Yoneda and Grossman (1995), summarized in Fig. 8, uses the thermodynamic data of Geiger et al. (1988) for grossite and hibonite and predicts a CHP sequence for condensation from a solar gas at pressures from $\sim 5 \times 10^{-3}$ (S. Yoneda, unpubl. data) through the lowest pressure modelled, 10-6 atm. Equilibrium condensation temperatures decrease with decreasing pressure. In the case of corundum, the temperature of the start of condensation, with no enrichment in dust relative to gas, ranges from 1770 K at 10^{-3} atm to 1571 K at 10^{-6} atm (Yoneda and Grossman, 1995). At pressures above the heavy dashed line in Fig. 8, corundum is not the first condensate according to these thermodynamic data.

The Geiger et al. (1988) data were preferred by Yoneda and Grossman (1995), but these data are not consistent with the liquid model of Berman (1983), which was used by Yoneda and Grossman (1995) for calculations in which liquid was predicted to be stable. Different data sets give different results. The Berman (1983) data for grossite and hibonite stabilize grossite relative to hibonite compared to the Geiger *et al.* (1988) data, predict grossite condensation before perovskite, and thus yield a corundum-hibonite-grossite sequence over much of the Geiger et al. (1988) CHP field in Fig. 8. As indicated by the horizontal lines in Fig. 8, the CHP sequence is restricted to pressures <10-4 atm and CI dust/gas enrichments <10× solar (for $P^{\text{tot}} \ge 10^{-6}$ atm) when the Berman (1983) data are used, while use of the Geiger et al. (1988) data expands the field for the CHP condensation sequence to a much wider range of pressures and dust/gas ratios (vertical lines only), especially between 10-3 and 10-6 atm, which are generally considered the most likely pressures at which condensation took place in the solar nebula. Both data sets indicate that the CHP condensation sequence is favored by low total pressure and low dust/gas enrichment. Except for CH chondrites (e.g., Grossman et al., 1988; Brearley and Jones, 1998), grossitebearing inclusions are very rare. Inclusions with hibonite and perovskite are much more common than hibonite-grossite



log CI dust/gas enrichment factor

FIG. 8. A summary of the variation of equilibrium condensation sequence with total pressure (P^{tot}) and CI chondritic dust/gas enrichment relative to solar composition. Plotted points represent conditions for which thermodynamic calculations have been performed using data from Geiger *et al.* (1988) for hibonite and grossite. The range of conditions for which a corundum–hibonite–perovskite (CHP) condensation sequence is predicted using these data is represented by vertical lines; when data for hibonite and grossite from Berman (1983) are used, the CHP sequence is confined to the region with horizontal lines. The two points labelled C-H-CA have a corundum–hibonite–CaAl₂O₄ sequence. The extent of conditions for which this sequence is predicted has not been explored.

inclusions, so it is possible that the thermodynamic data of Berman (1983) overestimate the stability of grossite. Meteoritic hibonite contains Mg and Ti, however, and the degree to which these substitutions increase the stability of hibonite is not accounted for by either data set. Despite this limitation, we can say that if the Berman (1983) data are correct, and if the corundum-hibonite-grossite inclusion reported by Krot *et al.* (2001) is a condensate, then it could have formed at higher P^{tot} and/or higher dust/gas enrichment than M98-8.

Corundum-Bearing Inclusions: Implications for Distribution of Aluminum-26 in the Early Solar Nebula

Corundum-bearing inclusions are very rare, probably because if they were not removed from the nebular gas soon after they formed, the corundum would have continued to react with the gas to form hibonite. In addition to M98-8 and GR-1, two other corundum-bearing samples have been analyzed by ion probe. They both may be condensates from the early solar nebula but do not have many features in common. Murchison sample BB-5 (Bar-Matthews *et al.*, 1982) has a corundum-rich core enclosed in hibonite in a texture that has small void spaces and is more compact than that of M98-8. Like M98-8, BB-5 has chondrite-normalized heavy REE abundances > light REE and no excess ²⁶Mg, indicating that the sample did not contain live ²⁶Al when it formed. Murray sample F5 has a group II REE pattern, and it contains excess ²⁶Mg consistent with *in situ* decay of ²⁶Al and an initial ²⁶Al/²⁷Al ratio of $(4.1 \pm 0.2) \times 10^{-5}$ (Fahey, 1988). Of the three corundum-bearing inclusions that may be primary condensates, two (BB-5 and M98-8) did not contain live ²⁶Al when they formed, although many other, less refractory (*e.g.*, melilite-bearing) CAIs did (MacPherson *et al.*, 1995).

Because evidence for ²⁶Al is found in many, but not all, refractory inclusions, the origin of this radionuclide and its significance for constraining the timescales of formation of CAIs are controversial. From its short half-life $(7.3 \times 10^5 \text{ years})$ we know that ²⁶Al must have entered CAIs soon (*i.e.*, not more than a few million years) after its synthesis. This relatively short timescale has led to two competing suggestions: either ²⁶Al formed in the solar system by irradiation of CAIs (or their precursors) by solar energetic particles, as in the X-wind model (e.g., Shu et al., 1997; Lee et al., 1998; Gounelle et al., 2001), or it was added as a spike of freshly synthesized nuclides from a single stellar source (e.g., Lee et al., 1977; Sahijpal and Goswami, 1998; Goswami et al., 2001). In its favor, the X-wind model provides a mechanism for removal of CAIs from their formation region, which is necessary for the preservation of high-temperature assemblages. Also, that there was some radionuclide production by solar particle irradiation is evident from the existence of live ¹⁰Be in CAIs (McKeegan et al., 2000), as all Be is destroyed in stellar interiors and ¹⁰Be can only be made by spallation, either in the interstellar medium, where the galactic cosmic ray flux is low, or by solar flares near the Sun. Serious problems with an analogous origin for ²⁶Al, and evidence for seeding of the solar nebula by radioactive stellar debris, however, result from the observation that in hibonite, ²⁶Al almost always occurs with an even shorterlived ($t_{1/2} \approx 10^5$ years) radionuclide, ⁴¹Ca (Sahijpal and Goswami, 1998; Sahijpal et al., 1998, 2000). Calculations using measured and calculated nuclear cross-sections show that particle irradiation cannot simultaneously produce the correct initial ²⁶Al and ⁴¹Ca abundances from the same target material (Shu et al., 1997; Sahijpal and Goswami, 1998; Lee et al., 1998; Sahijpal et al., 2000; Goswami et al., 2001).

This problem led proponents of irradiation (Shu *et al.*, 1997, 2001; Gounelle *et al.*, 2001) to suggest that ²⁶Al was made in thick (~3 mm), Ca-free ferromagnesian mantles that enclosed CAIs and provided sufficient shielding to yield the observed initial ²⁶Al without overproduction of ⁴¹Ca. The models for mantle formation, however, ignore major petrologic constraints. Shu *et al.* (2001) invoke formation of the mantles by melting of chondritic dust and generation of immiscible liquids, but experimentally determined phase equilibria show that chondritic melts do not give rise to immiscible liquids. Gounelle *et al.*

(2001) likened the mantle-forming process to the "zone refining" technique used for purifying metal. For this to work on proto-CAIs, a heat pulse would have to move from the outside of the CAIs inward, driving the residual melt toward the cores, but during heating by solar flares, as envisioned by Gounelle *et al.* (2001), the insides of the objects will be no hotter than the outsides. This would yield objects with forsteritic interiors and Ca-, Al-, Si-rich exteriors containing no corundum, hibonite or melilite, because partial melting of chondritic matter cannot produce a residue of CAI composition. The accretionary rims that are observed around CAIs do not meet the requirements of the X-wind model. The rims reach 3 mm in thickness only where they fill embayments in the host CAIs, and they are not Ca-free (MacPherson *et al.*, 1985).

Even if suitable ferromagnesian mantles did exist, it would be very difficult, perhaps impossible, to transport ²⁶Al from them into CAIs in a way that yields not only a dominant initial ²⁶Al/²⁷Al ratio among many CAIs, but also accounts for the fact that phases in CAIs fall on Al-Mg isochrons. The existence of isochrons requires that the ²⁶Al/²⁷Al ratio was uniform throughout each CAI as an initial condition, further requiring that, when ²⁶Al was incorporated, the constituent minerals of such CAIs equilibrated with a common fluid phase, probably implying that the CAIs were molten. The Gounelle et al. (2001) model calls for entry of ²⁶Al into the nebular gas by vaporization of the ²⁶Al-bearing ferromagnesian mantles without vaporization of the CAIs themselves. Actually, the ²⁶Al would probably not evaporate, but would stay in the proto-CAIs even as they underwent the nearly complete melting implied by the isochrons. Melting of the entire proto-CAI + mantle assemblage after ²⁶Al production would not be allowed, because this would result in a homogeneous droplet much more Mg- and Si-rich than observed CAIs. Formation of CAIs as evaporative residues of compositions this rich (i.e., near-chondritic) in Mg and Si would produce much larger Mg and Si isotopic massfractionations than are observed in CAIs. It is highly unlikely, however, that the ferromagnesian mantles could be heated and evaporated without some dissolution of them into the molten CAI cores, also requiring more extensive Mg and Si evaporation and isotopic fractionation than are indicated by the isotopic compositions of CAIs. Finally, in another unlikely scenario, at very particular temperatures at low nebular pressures, solid-tovapor evaporation of mantles could occur while solid CAI cores remain intact. A subsequent melting event would then be required to permit incorporation of ²⁶Al uniformly into the CAIs.

In light of the problems of coproduction of the observed abundances of radionuclides by energetic particle irradiation, arguments for a stellar source for ²⁶Al and ⁴¹Ca are discussed by Sahijpal and Goswami (1998) and Sahijpal *et al.* (1998, 2000). Models must account for the uniformity of initial abundances of ²⁶Al and ⁴¹Ca and for the formation of hibonite with both and without both of these radionuclides. Three possibilities for the formation of ²⁶Al-free inclusions, such as M98-8, were considered by Sahijpal and Goswami (1998) and

Sahijpal et al. (2000). They are (1) formation before injection of ²⁶Al into the nebula was completed; (2) formation after ²⁶Al decay; and (3) heterogeneous distribution of ²⁶Al in the solar nebula. Their analyses and data from the literature (e.g., Clayton et al., 1988; MacPherson et al., 1995) show that inclusions that did not incorporate ²⁶Al also did not incorporate ⁴¹Ca and exhibit large isotopic anomalies of nucleosynthetic origin in ⁴⁸Ca and ⁵⁰Ti relative to most CAIs. They interpreted these features as primary nucleosynthetic signatures. In rejecting the heterogeneous distribution model, these workers noted the conclusion by MacPherson et al. (1995) that Al isotopic heterogeneity appears to have been minor among CAI precursors, which were strongly dominated by materials with initial $^{26}Al/^{27}Al$ ratios of 5 ×10–5. This interpretation of the data is consistent with the nebular evolution models of Foster and Boss (1997), which predict limited heterogeneity of ²⁶Al due to steady (as opposed to episodic) injection of live radionuclides during the collapse of the solar nebula. Formation of ²⁶Al-free inclusions after decay of the radionuclide is also not favored, because this would require formation of ²⁶Al-bearing inclusions early, storage of these inclusions in the nebula for several million years, then formation of even more highly refractory inclusions, and incorporation of ²⁶Al-free and ²⁶Al-bearing inclusions into the same parent bodies. Assuming a stellar source for the radionuclides and that the travel time between the source and the protosolar cloud was negligible, Sahijpal and Goswami (1998) and Sahijpal et al. (2000) concluded that ²⁶Al-free hibonite grains and hibonite-, corundum-rich refractory inclusions formed very early in the history of the solar system, during a presumed about $(2-6) \times 10^5$ year interval between impact of the stellar shock front that triggered the collapse of the solar nebula and the injection of live radionuclides (26Al and 41Ca) into the CAI-forming region. This model has the advantages of (a) explaining the correlation between ²⁶Al and ⁴¹Ca abundances in CAIs and (b) not invoking a several million-year gap between generations of CAIs.

The present sample, M98-8, did not contain live ²⁶Al when it formed, and within the context of the Sahijpal *et al.* (2000) model it would therefore correspond to a condensate assemblage from the earliest phases of the solar nebula. Sample F5 is a hibonite-rich sample that did contain ²⁶Al when it formed, and it has a group II REE pattern (Fahey, 1988), which is indicative of prior volatility-related fractionation of its precursors. This suggests that its formation occurred somewhat later.

Formation of Oxygen-16-Rich Inclusions

Ion probe analyses show that the most refractory phases of CAIs tend to be the most ¹⁶O-rich, with $\delta^{18}O \approx \delta^{17}O$ and typically between -40 and -50‰ (*e.g.*, Ireland *et al.*, 1992). These are thought to represent the primary isotopic compositions of the phases, and therefore that of the early nebular vapor, while phases with isotopic compositions closer to normal are thought to have obtained their compositions

through reaction with a relatively ¹⁶O-poor gas (Clayton et al., 1977). Although ¹⁶O is made in supernovae, as is ²⁶Al, the ¹⁶O enrichments are not correlated with ²⁶Al and ⁴¹Ca contents. Hibonite tends to be 16 O-rich, about -50%, whether or not it contained live ²⁶Al when it formed (Fahey et al., 1987). This implies that ¹⁶O was not injected into the nebula along with the short-lived nuclides. It is not known how the nebular gas became ¹⁶O-rich. Some workers favor evaporation of ¹⁶O-rich dust to form ¹⁶O-rich gas (Scott and Krot, 2001; Cassen, 2001). Their models, based on observed compositions of phases in CAIs and ¹⁶O-rich endmembers assumed to have $\delta^{17}O = -50\%$, suggest that ¹⁶O-rich phases in CAIs formed from a vapor enriched in dust by a factor of ~30 relative to solar composition. A corundum-hibonite-perovskite condensation sequence would still be possible at this degree of dust enrichment at low Ptot (Fig. 8), but for $P^{\text{tot}} \approx 10^{-3}$ atm or higher, corundum would not be the first condensate and condensation of liquid would be likely (Yoneda and Grossman, 1995; Ebel and Grossman, 2000). If dust enrichment is required to produce ¹⁶O-rich minerals, this would place an upper limit on the range of pressures at which inclusions thought to be gas-to-solid condensates could have formed. Another way to produce an ¹⁶O-rich gas may be by non-mass-dependent isotopic fractionation during gas-phase chemical reactions (e.g., Thiemens, 1999), but this process has not been shown to work for either SiO or CO, the dominant oxygen-containing molecules in a solar gas.

CONCLUSIONS

M98-8 consists of corundum, hibonite, and perovskite, the first three condensate minerals predicted to form by equilibrium condensation from the solar nebula under certain conditions. From petrographic observations, chemical and isotopic data, and the inability of any model that includes a molten stage, relic grains or evaporation to account for the features of this inclusion, we conclude that it is a primary condensate from an ¹⁶O-rich gas in the early solar nebula. Thermodynamic condensation calculations show that a corundum-hiboniteperovskite sequence is possible under a fairly wide range of total pressures $\leq 5 \times 10^{-3}$ atm, including those typically favored for refractory inclusion formation. The lower the pressure, the higher the permissible dust/gas enrichment, up to between 100 and $1000 \times \text{solar}$ for $P^{\text{tot}} = 10^{-6}$ atm. The sample did not contain live ²⁶Al when it formed, which is consistent with the Sahijpal and Goswami (1998) model for the injection of ²⁶Al into the solar nebula, which suggests that ²⁶Al-free refractory inclusions are among the first solids formed in the solar system. If this model is correct, then M98-8 formed within $\sim 6 \times 10^5$ years of the beginning of the collapse of the protosolar cloud.

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