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# A short timescale for changing oxygen fugacity in the solar nebula revealed by high-resolution <sup>26</sup>Al-<sup>26</sup>Mg dating of CAI rims

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#### **Abstract**

Most rocky objects in the solar system, including the primitive chondrites and the terrestrial planets themselves, formed at oxygen fugacities ( $f_{\rm O2}$ ) near that of the Iron–Wüstite (IW)  $f_{\rm O2}$  buffer. Conversely, the most ancient rocky objects of the solar system, the calcium aluminum-rich inclusions (CAIs), formed at  $f_{\rm O2}$  values 5 orders of magnitude lower than the IW buffer in an environment more closely resembling a solar gas. High-resolution Mg isotope data and estimates for  $f_{\rm O2}$  for rims on CAIs show that this shift from ~solar to protoplanetary (chondritic)  $f_{\rm O2}$  occurred in 100,000 to 300,000 yr for these objects. Magnesium isotopes show further that the rise in  $f_{\rm O2}$  was accompanied by a rise in the partial pressure of Mg. These results establish that CAIs entered a region resembling where planet progenitors formed within  $3 \times 10^5$  yr of their formation in the solar nebula.

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# 1. Introduction

Refractory calcium aluminum-rich inclusions (CAIs) represent the most primitive record of rock formation in the solar system. Astrophysical models

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for the evolution of solids in the young solar protoplanetary disk must account for this record. Young protoplanetary disks in general evolve by viscous accretion to the star coupled with outward transport of angular momentum; as a consequence of viscous behavior, some disk materials spiral inwards towards the growing star while others move outward. How CAIs fit into this basic evolutionary picture of disk evolution is poorly understood. In this study we use

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so-called Wark-Lovering rims on CAIs as a record of time-dependent changes in conditions in the solar protoplanetary disk.

Wark-Lovering (WL) rims [1] makeup the outermost portion of many CAIs. They consist of concentric bands tens of micrometers (µm) thick. Each band is composed of a distinctive mineralogy and they are often monomineralic. The bands comprise a characteristic mineralogical sequence that is independent of differences among CAI interiors. The normal sequence moving from the interior towards the edge of the CAI is spinel  $\pm$  perovskite  $\pm$  hibonite,  $\pm$  melilite, Al-rich clinopyroxene, and  $\pm$  forsterite. These enigmatic features have been studied for decades. Some workers suggest that the rims formed by condensation. Others conclude that they were evaporative residues. Their significance has proven elusive even as their ubiquity betrays their importance as a consequence of a process fundamental to the early solar nebula [1]. Here we use the amount of radiogenic <sup>26</sup>Mg (<sup>26</sup>Mg\*) due to decay of the extinct radionuclide <sup>26</sup>Al (mean life=1.05 million yr) to date WL rim formation relative to CAI interiors. At the same time we apply measured Ti<sup>3+</sup>/Ti<sup>4+</sup> and <sup>25</sup>Mg/<sup>24</sup>Mg as "cosmobarometers" to constrain the  $f_{\rm O2}$  (oxygen fugacity) and  $P_{\rm Mg}$  (partial pressure of Mg) attending growth of the CAIs and their rims. The picture that emerges is one of rapidly changing gas-phase conditions in the early solar system as recorded by the CAIs. This timescale can be compared with those for protoplanetary disk evolution in general.

The growth model for WL rim formation described herein is similar to the metasomatic model [2] in so far as both rely on large gradients in chemical potentials to drive the formation of the WL rims. In addition, both models invoke reaction between the CAI and a surrounding Mg and Si-rich gas. However, a metasomatic mechanism implies that refractory elements are scavenged from the interior while we show that growth could have occurred from elements in the gas phase alone. We show further that the new Mg isotope data for WL rims are more easily understood if the rims represent the product of growth that is chemically and isotopically independent from the host CAIs. Growth in response to high chemical potential gradients can explain both the uncoupling between interior and rim formation and the WL rim and host CAI Mg isotope data.

# 2. Analytical methods

# 2.1. Magnesium isotopes

Measurements of <sup>25</sup>Mg/<sup>24</sup>Mg, <sup>26</sup>Mg/<sup>24</sup>Mg and <sup>27</sup>Al/<sup>24</sup>Mg in the CAI interiors and rims were obtained by ultraviolet (UV) laser ablation combined with multiple-collector inductively coupled plasmasource mass spectrometry (MC-ICPMS) using methods described previously [3]. The laser was operated at a 213 nm fluence of 20 to 30 J/cm<sup>2</sup> and a pulse repetition rate of 1 to 2 Hz. Analytical spot size was varied between 50 and 100 µm depending upon Mg concentration. Rims were analyzed using 50 µm wide trenches measuring ~200 µm in length. Excess radiogenic  $^{26}$ Mg values were calculated from measured  $^{26}$ Mg/ $^{24}$ Mg and  $^{25}$ Mg/ $^{24}$ Mg expressed in the linear delta notation ( $\delta'$ ) relative to the DSM3 magnesium standard [4] where  $\delta^{i}Mg' = \ln(({}^{i}Mg/{}^{24}Mg)_{sample}/({}^{i}Mg/{}^{24}Mg)_{DSM3})10^{3}$  and i represents either 25 or 26. On the DSM3 scale,  $\delta^{25}$ Mg' of typical chondrite material (e.g., bulk CI meteorite) is 0.0 %. With these definitions for  $\delta'$  values, radiogenic <sup>26</sup>Mg is quantified as  $\delta^{26} Mg^* = \delta^{26} Mg' - \delta^{25} Mg' / (0.521)$  where 0.521 is the slope of the equilibrium mass-dependent isotope fractionation relationship between  $\delta^{26}$ Mg' and  $\delta^{25}$ Mg'( $\beta$ ). The exact value of the mass-dependent fractionation relationship depends upon physicochemical process and ranges from 0.521 to 0.511 [5]. Our conclusions are not altered by the choice of  $\beta$  value within this range because WL rims have  $\delta^i$ Mg' values near zero. The use of multiple detectors and the sample-standard comparison method for correcting for instrumental mass bias affords routine analysis of samples that have <sup>27</sup>Al/<sup>24</sup>Mg values less than five such as the WL rims [6]. Tests with solids and solutions show that within realistic elemental concentrations for CAIs matrix mass bias effects are well within the reproducibility (0.25%/amu, 2 s.d.) of our solid-solid standard comparison method (cf. [6]). External reproducibility is estimated from repeated analyses of USNM 136718 forsterite, Burma spinel, and a synthetic fassaite glass (glass P10). A <sup>48</sup>Ca<sup>++</sup> interference on m/z=24 produces a consistent enrichment of  $\sim 0.04\%$  in  $\delta^{26}$ Mg\* per unit Ca/Mg. A correction for the interference was applied based on the Ca/Mg ratios of the analyzed materials as determined from their measured Al/Mg ratios and chemical formulae (e.g., in melilite Ca/Mg=2/(1-x) where x is the mole fraction of the tschermak exchange component  $Al_2Mg_{-1}Si_{-1}$  in the melilite). Tests with terrestrial melilites, vesuvianite, and glass P10 demonstrate that this correction scheme affords accurate results within the precision of the measurements.

# 2.2. Oxygen fugacity based on $Ti^{3+}/Ti^{4+}$ in pyroxene

The  $f_{\rm O2}$  values attending pyroxene growth in the interiors and the rims of the CAIs were determined from the ratio of the activity of CaTi<sup>4+</sup>Al<sub>2</sub>O<sub>6</sub> to the activity of CaTi3+AlSiO6 in pyroxene. This activity ratio defines  $f_{\rm O2}$  by virtue of gas-solid equilibria involving O2. The oxidation state of Ti was obtained by calculation of relative abundances of cations from electron microprobe analyses (UCLA Jeol superprobe) assuming no vacancies in the pyroxene formula. This approach has been shown to be robust for CAI pyroxenes because of the virtual absence of other multivalent cations (e.g., Fe) [7,8]. Twenty nine analyses of pyroxene in the WL rim of CAI 144A were obtained using a minimum electron beam size. The TiO<sub>2</sub> content (total Ti as TiO<sub>2</sub>) of the rim pyroxenes varied between 2 and 7 wt.% with most analyses being above 4 wt.%. Twenty three analyses representing 3 different Ti-rich pyroxenes located in the interior of CAI 144A were obtained for comparison with the rims. Interior pyroxene contained 17–19 wt.% TiO<sub>2</sub>. The amount of Ti<sup>3+</sup> was calculated from total Ti based on oxygen excess in the cation-normalized formula (or cation deficiencies in the oxygen normalized formula). Reported uncertainties in the final cation assignments are based on a Monte Carlo error simulation (n = 300 trials) in which the uncertainties in the wt.% oxide data were propagated through the cation norm calculation. The algorithm for the integrated probability density function for a Gaussian distribution used to model uncertainties is described by Harbaugh and Bonham-Carter [9]. Random numbers generated to draw from parent populations for each oxide come from an algorithm described by Press et al. [10].

# 3. Calcium Aluminum-rich inclusions

Rims and interiors of three igneous CAIs from the Leoville and Allende CV3 carbonaceous chondritic meteorites were examined as part of this study. Allende 3576-1 "b" is a  $\sim 4 \times 3.5$  mm Type B CAI composed of intergrown melilite and pyroxene surrounded by a mantle dominated by melilite. Spinel is abundant throughout the inclusion. A WL rim that is ~30-60 µm thick and composed of spinel and pyroxene coats the melilite dominated interior. Leoville 144A is a compact, oval ~10 × 6 mm Type A CAI composed of fine grained intergrown melilite and rare Ti-Al-rich pyroxene that encloses abundant micronsized perovskite grains. Spinel is distributed throughout the CAI. The WL rim of Leoville 144A is composed of an inner layer of Mg-rich spinel intergrown with platy crystals of hibonite, thin (<10 μm) discontinuous layers of melilite and calcic pyroxene and an outer layer composed predominantly of Al-rich pyroxene. The spinel in the inner layer encloses numerous small perovskite grains. The thickness of the outer pyroxene layer as presented in the polished surface used for these analyses varies from <50 to ~200 μm. Leoville MRS3 is a fragment of a Type A CAI measuring ~0.7 mm in length and 0.5 mm in width. It is largely composed of coarse grained melilite with subordinate spinel. The WL rim of MRS3 is <50–100 um thick and is composed of an inner layer of spinel and an outer layer of Al-rich pyroxene.

# 4. Magnesium isotope ratios in Wark-Lovering rims and interiors of CAIs

# 4.1. <sup>26</sup>Al as a chronometer

The one-time presence of  $^{26}$ Al in the solar system is evidenced by excesses of its decay product,  $^{26}$ Mg\* ( $^{26}$ Al decays to radiogenic  $^{26}$ Mg,  $^{26}$ Mg\*, primarily by  $\beta^+$  emission and also by electron capture), correlated with Al/Mg in the constituents of meteorites [11]. The use of  $^{26}$ Al as a chronometer relies on variations in the initial  $^{26}$ Al/ $^{27}$ Al, hereafter ( $^{26}$ Al/ $^{27}$ Al)0, in objects formed within several mean lives of  $^{26}$ Al decay. Values for ( $^{26}$ Al/ $^{27}$ Al)0 are defined by isochrons comprising linear correlations between  $^{26}$ Mg\*/ $^{24}$ Mg (expressed as  $\delta^{26}$ Mg\*, the per mil deviation in  $^{26}$ Mg/ $^{24}$ Mg from the value corresponding to mass-dependent isotope fractionation) and  $^{27}$ Al/ $^{24}$ Mg. The slopes of these isochrons are numerically equivalent to ( $^{26}$ Al/ $^{27}$ Al)0 since all of the  $^{26}$ Al decayed away

billions of years ago. Age differences are reflected in differences in  $(^{26}\text{Al}/^{27}\text{Al})_0$  if the initial  $^{26}\text{Al}/^{27}\text{Al}$  was uniform in the early solar system. Recent UV laser ablation and acid digestion MC-ICPMS analyses of CAIs [3,6,12,13] show that the  $(^{26}\text{Al}/^{27}\text{Al})_0$  for CAIs in the early solar system was at least  $6\times 10^{-5}$  and possibly as high as  $7\times 10^{-5}$  (but see Bizzarro et al. [14]), and that the canonical  $(^{26}\text{Al}/^{27}\text{Al})_0$  of  $4.5\times 10^{-5}$  is a value representing the time at which heating of CAIs ceased.

Our UV-MC-ICPMS data show that the WL rims formed within ~300,000 yr of the first CAI interiors (Table 1). Measurements of WL rims from Leoville 144A, Leoville MRS3 and Allende 3576-1 "b" define an Al–Mg isochron corresponding to an  $(^{26}\text{Al}/^{27}\text{Al})_0$  value of 5.3  $(\pm\,0.8)\times10^{-5}$  with a  $\delta^{26}\text{Mg*}$  intercept of zero  $(0.06\pm0.1)$  (Fig. 1). Interiors of the Leoville CAIs display well-defined  $(^{26}\text{Al}/^{27}\text{Al})_0$  of at least  $6\times10^{-5}$  [3] (Appendix A, an electronic supplement). The data for the Allende inclusion are less well behaved (apparently due to alteration) but are consistent with the Leoville results. From the difference between the rim  $(^{26}\text{Al}/^{27}\text{Al})_0$  value and those representing growth of the CAI interiors the time interval between initial CAI growth and

rim formation can be constrained from the expression  $(5.3 \times 10^{-5})/(6.0 \times 10^{-5}) = \exp(-\lambda \Delta t)$ . The  $\Delta t$  obtained is 130,000 yr based on a decay constant  $\lambda$  of  $9.52 \times 10^{-7}$  yr<sup>-1</sup>. This value may be a minimum. The  $(^{26}\text{Al}/^{27}\text{Al})_0$  of the interior of Leoville 144A could have been as high as  $7 \times 10^{-5}$  meaning that  $\Delta t$  could have been as large as 290,000 yr. Alternatively, if  $^{26}\text{Al}$  is heterogeneous in the early solar system, then the time difference is likely to be even smaller.

# 4.2. <sup>25</sup>Mg/<sup>24</sup>Mg as a cosmobarometer

CAI interiors and rims have markedly different  $\delta^{25} \mathrm{Mg'}$  suggestive of distinctly different  $P_{\mathrm{Mg}}$  during their formation. Interiors of the three igneous CAIs have  $\delta^{25} \mathrm{Mg'}$  values significantly greater than the 0% ( $\pm$ 1) that characterizes Earth, Moon, and most constituents of chondritic meteorites [5,15]. The WL rims, on the other hand, have  $\delta^{25} \mathrm{Mg'} \leq 0\%$  irrespective of the  $\delta^{25} \mathrm{Mg'}$  values of the enriched interiors (Fig. 2).

The depleted  $\delta^{25} \text{Mg}'$  compositions observed in the WL rims indicate that they are condensates formed at relatively high  $P_{\text{Mg}}$  while the high  $\delta^{25} \text{Mg}'$  values of

Laser ablation Mg isotope data for Wark–Lovering rims on CAIs from CV carbonaceous chondrites

Analysis	Phase(s)	Location from rim (µm)	$^{27}Al/^{24}Mg$	$\sigma_{\mathrm{m}}$	$\delta^{25}$ Mg'	$\sigma_{ m m}$	$\delta^{26}$ Mg'	$\sigma_{ m m}$	$\delta^{26}$ Mg*	$\sigma_{ m m}$
Allende (3.	576-1" b" )									
Spot 20	W-L>>mel	-50	0.57	0.03	-0.86	0.19	-1.53	0.12	0.12	0.32
Spot 21	W-L>>mel	-50	0.24	0.02	-1.30	0.15	-1.81	0.14	0.69	0.19
Leoville (1	<i>44A)</i>									
Line 3	W-L Al-Ti diopside	-100	0.76	0.04	-0.09	0.12	0.11	0.06	0.27	0.23
Line 4	W-L Al-Ti diopside	-75	1.04	0.09	-1.74	0.11	-2.74	0.11	0.60	0.17
Line 5	W-L Al-Ti diopside	-150	0.94	0.01	-0.38	0.17	-0.48	0.13	0.25	0.28
Line 6	W-L hb sp	-50	0.91	0.04	-0.85	0.11	-1.33	0.12	0.30	0.19
Line 7	W-L Al-Ti diopside	-120	0.96	0.02	-1.29	0.09	-2.32	0.19	0.16	0.15
Line 8	W–L hb sp	-25	1.31	0.07	-1.97	0.14	-3.39	0.10	0.39	0.29
Leoville (M	MRS3)									
Line 1	W-L Al-Ti diopside	-20	4.62	0.54	-0.45	0.12	0.81	0.26	1.67	0.32
Line 2	W-L Al-Ti diopside	0	9.02	0.70	-0.98	0.31	2.09	0.38	3.97	0.49

Measurements are a mixture of WL rim phases; dominant phase is indicated when known. Excess radiogenic  $^{26}$ Mg values were calculated from measured  $^{26}$ Mg/ $^{24}$ Mg and  $^{25}$ Mg/ $^{24}$ Mg expressed in the linear delta notation ( $\delta'$ ) relative to the DSM3 magnesium standard where  $\delta^i$  Mg'=ln(( $^i$ Mg/ $^{24}$ Mg)<sub>sample</sub>/( $^i$ Mg/ $^{24}$ Mg)<sub>DSM3</sub>)10<sup>3</sup> and i represents either 25 or 26 (CI chondrite=0.0‰ on the DSM3 scale [5]). With these definitions for  $\delta'$  values radiogenic  $^{26}$ Mg is  $\delta^{26}$ Mg\*= $\delta^{26}$ Mg' -  $\delta^{25}$  Mg'/(0.521) where 0.521 is the slope of the equilibrium mass-dependent isotope fractionation relationship between  $\delta^{26}$ Mg' and  $\delta^{25}$ Mg'. Interior Mg isotope measurements can be found in the electronic supplement.

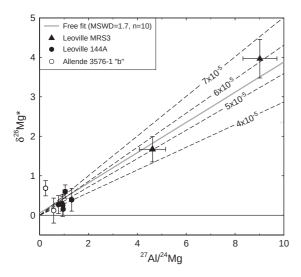


Fig. 1. The Al–Mg evolution of WL rims as defined by those measured from Leoville 144A, Leoville MRS3, and Allende 3576-1 "b" (best fit for initial  $^{26}$ Al/ $^{27}$ Al=5.3×10 $^{-5}$ ±0.8, 1 $\sigma_{\rm m}$ , MSWD=1.7, n=10). Dashed lines for reference  $^{26}$ Al/ $^{27}$ Al values as indicated. The Mg data for WL rims were obtained in situ by UV laser ablation MC-ICPMS, reported in Table 1. Measurements were obtained as ~50 μm diameter spots and ~50 μm wide line scans. Reported uncertainties are 1 $\sigma_{\rm m}$ .

the interiors of these igneous CAIs (the igneous CAIs were once molten in space) shows that they are evaporative residues formed at lower  $P_{\rm Mg}$  [16] (Fig. 2). The role of Mg isotope ratios as Mg barometers is seen with reference to an equation that governs the net flux of Mg volatilized from a molten sphere [17], rewritten in the form [5]:

$$J_{\rm Mg,net} = \frac{J_{\rm Mg,evap} \left( 1 - \frac{P_{\rm Mg,\infty}}{P_{\rm Mg,sat}} \right)}{1 + \Gamma} \tag{1}$$

in which,

$$\Gamma = \frac{\gamma_{\rm Mg} r}{D_{\rm Mg,gas}} \sqrt{\frac{RT}{2\pi m_{\rm Mg}}} \tag{2}$$

and where  $J_{\rm Mg,net}$  is the net difference between the evaporative and condensation fluxes for Mg (Mg evaporates as Mg gas),  $J_{\rm Mg,evap}$  is the evaporative flux of Mg,  $P_{\rm Mg,\infty}$  is the partial pressure of Mg far removed from the molten object,  $P_{\rm Mg,sat}$  is the saturation partial pressure of Mg,  $D_{\rm Mg,gas}$  is the diffusion coefficient of Mg through the gas phase,  $\gamma_{\rm Mg}$  is the evaporation factor for Mg, r is the radius of the molten object,

and  $m_{\rm Mg}$  is the mass of the volatilizing species. In terms of this equation, isotope fractionation will occur when  $J_{\rm Mg,net}/J_{\rm Mg,evap} \rightarrow 1$  while no fractionation occurs when  $J_{\rm Mg,net}/J_{\rm Mg,\ evap} \rightarrow 0$ . The latter occurs where the background pressure  $P_{\rm Mg,\ sat}$ . This could be the case where a population of volatilizing liquid spheres contributes to an elevated background of partial pressure of rock-forming elements by virtue of a relatively high number density. In the other extreme where  $P_{\rm Mg,\ sat} >> P_{\rm Mg,\ \infty}$ , as would be the case where objects

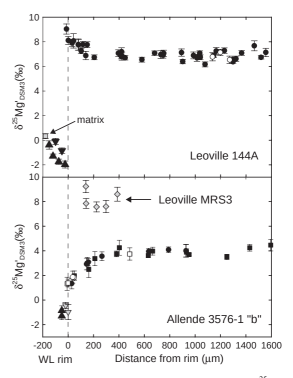


Fig. 2. Systematic variations in intrinsic Mg isotope ( $\delta^{25}$ Mg') composition obtained by in situ laser ablation core-to-rim traverses of studied CAIs. All CAIs possess WL rims that have low  $\delta^{25}$ Mg' compositions relative to chondritic, but exhibit interiors with  $\delta^{25}$ Mg' higher than chondritic values by ~3% to>6%. Closed symbols are melilite, open symbols are pyroxene, and triangles are WL rims, unless labeled otherwise. (Top) Six radial traverses of Leoville 144A exhibit a progressively enriched edge (<200  $\mu$ m), a flat bottom interior, and a reproducible subchondritic WL rim that exhibits outward per mil enrichment converging on chondrite. (Bottom) Seven analyses of Leoville MRS3 are consistent with those of Leoville 144A and three radial traverses of Allende 3675-1 "b" yield a progressively depleted margin (~500  $\mu$ m), a flat topped interior and a subchondritic WL rim. Reported uncertainties are  $2\sigma_{\rm m}$ .

are not sufficiently close together to cause a pervasive rise in background partial pressures, the only way to prevent fractionation is for  $\Gamma$ , and therefore total gas pressure, to be large (e.g., [12]). The kinetic theory of gases permits one to express  $\Gamma$  as a function of pressure. Such calculations show that  $\Gamma$  is large enough to preclude fractionation when total pressures approach 1 bar (e.g., using experimental values for  $\gamma_i$ =0.06 [18],  $\Gamma$ =~10<sup>-2</sup> at 10<sup>-3</sup> bar and ~10 at 1 bar for a millimeter-sized object). From these considerations it is clear that the level of isotopic fractionation for a major rock-forming element like Mg in an object that was once molten in the early solar system is seen to be a barometer of either total pressure (large  $\Gamma$ ) or partial pressures relative to saturation (small  $\Gamma$ ).

Eqs. (1) and (2) show that high  $\delta^{25} \text{Mg}'$  in the interior of the once molten CAIs is a consequence of evaporation where  $P_{\text{Mg}}$  and total pressure were both low, and that the lack of Mg isotope fractionation among virtually all other rocky objects, including the WL rims and the millimeter-sized chondrules that were molten and free-floating in the nebula, signifies  $P_{Mg}$  values approaching saturation or high total pressures. Since the pressures required to raise  $\Gamma$  to sufficiently high values to preclude fractionation are several orders of magnitude greater than estimates for the nebula ( $\sim 10^{-3}$  bar maximum, e.g., [19]),  $P_{\text{Mg}}$  approaching saturation ( $P_{\text{Mg,sat}}$ ) is indicated.

# 4.3. Wark-Lovering rims are nebular condensates

Wark-Lovering rims have chondritic and subchondritic  $\delta^{25}$ Mg' values signifying that they, like the chondules and unlike CAIs interiors, formed at partial pressures of Mg approaching saturation. This in turns suggests that they formed by condensation. Wark and Lovering originally attributed the rims to nebular condensation, but after finding enrichments of rare earth elements (REE) and other refractory trace elements in rims, Wark and Boynton argued that rims were residues of intense (>2500 K) and brief (<2 s) thermal events that evaporated surface material from CAIs [20] that were then subsequently altered by a metasomatic process where inter-diffusion with later accreted olivine grains occurred (i.e., [2]). Wark and Boynton correctly recognized that spinel and Ti-Alpyroxene in the rims exhibit the same range of <sup>16</sup>Orich compositions as the interiors, and have therefore

not exchanged with, or been diluted by, greater amounts of "normal" oxygen (e.g., [21,22]). A similar, but more compelling argument can be made from Mg based on the fact that rims have excess <sup>26</sup>Mg. They also have even lower  $\delta^{25}$ Mg' than the surrounding matrix. Evaporation followed by accretion does not explain the low  $\delta^{25}$ Mg' of the rims. The enrichments in REE that are sometimes observed in the rims can be explained by the greater abundance of minerals that retain REE (primarily perovskite) stabilized by the condensation reactions. The present study suggests that the factor that stabilized pervasive perovskite in the rims may have been oxidation of Ti from Ti<sup>3+</sup> to Ti<sup>4+</sup>. In short, our high-precision Mg isotope data show that the WL rims are condensates and not evaporative residues.

# 5. A general model for Wark-Lovering rim formation

Ruzicka [2] and MacPherson and others [23] recognized the importance of large gradients in chemical potentials ( $\mu_i$ ) as the principal driving force for WL rim formation. Both studies emphasized the role of metasomatism (i.e., Si and Mg diffusion into the CAI). Here we consider that the isotope and  $f_{\rm O2}$  data for rims are consistent with new mineral growth and present thermodynamic calculations that reproduce the WL rim sequence as products of growth in high  $\mu_i$  gradients.

Thermodynamic calculations show that the characteristic sequence in WL rim mineralogy (spinel ± hibonite → Al-rich diopside → forsterite) is explained by placing solid melilite-rich CAI interiors at elevated temperature ( $\geq 1400 \text{ K}$ ) into a region that imposes high positive gradients in  $\mu Mg$  ( $P_{Mg}$ ) (and  $\mu SiO$  $(P_{SiO})$ ) and  $\mu O_2$   $(f_{O2})$ . The effects of high  $\mu Mg$  and  $f_{O2}$  can be seen in a phase diagram that shows the stability fields of minerals in the model system Ca-Mg-Al-Si-O as a function of the partial pressures of the gas species Mg, Ca, SiO, and O<sub>2</sub> (Fig. 3). For the purposes of illustration, Fig. 3 shows a section at fixed  $P_{SiO}$ . Similar diagrams can be constructed in which  $P_{SiO}$  is a variable. Partial pressures in Fig. 3 refer to a total standard state pressure of 1 bar, i.e., the  $P_i$ values are mole fractions at any total pressure. Solid lines delimit stability fields for anorthite, spinel,

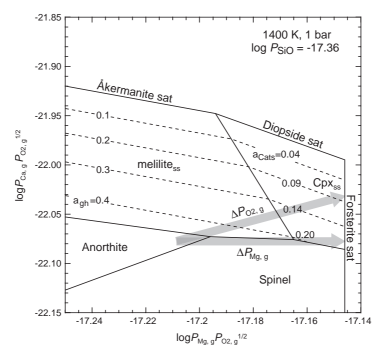


Fig. 3. Phase diagram of mineral stability in the model system Ca-Mg-Al-Si-O. This activity-activity diagram illustrates mineral growth expected during interaction of a CAI with an infinite reservoir of forsterite (i.e., chondritic)-equilibrated gas. Arrows indicates the sequence of mineral banding that will develop by evaporation-precipitation reactions where  $\log P_{\rm Mg, g}$  and  $\log P_{\rm O2, g}$  increase, respectively. Phase relations derived at 1400 K, 1 bar total pressure, using tabulated thermodynamic data [10,32], standard states of unit activity of pure minerals at all P and T and ideal gases at 1 bar and T, and an ideal-mixing-on-sites approximation for activities of solid solution components.

melilite and clinopyroxene. Lines defining the Alfree end members åkermanite, diopside, and forsterite represent saturation surfaces (solid lines). Gases with partial pressure (mole fractions) products higher than these limiting lines are metastable (supersaturated) with respect to these solid phases. Dashed lines represent isopleths of gehlenite and Ca-tschermak activities in melilite and clinopyroxene, respectively.

The model in Fig. 3 shows that a rise in  $P_{\rm Mg}$  and  $P_{\rm O2}$  ( $f_{\rm O2}$ ) at fixed  $P_{\rm Ca}$  (fixed partial pressure of Ca is expected since Ca is refractory and evaporation of rock dust would result in relatively negligible transfer of Ca to the gas phase) will produce the general sequence of mineral zones found in WL rims. Changes in temperatures and/or partial pressures of SiO, Mg, Ca, and/or  $O_2$  along the reaction path would produce variations in the rim mineralogy. For example, polythermal growth of the rims is a satisfactory explanation for the presence or absence of a significant melilite layer between spinel and Al-rich pyrox-

ene because the melilite stability field will expand or contract substantially with changes in temperature. Moreover, initially higher temperatures can explain the origin of hibonite in the inner layer.

In any case, the phase diagram predicts that a rise in the partial pressures of the major chondritic rockforming elements Mg and O will lead to mineralogies resembling WL rims. The low  $\delta^{25}$ Mg' of the WL rims relative to their host CAIs is an indication that  $P_{\rm Mg}$  was indeed relatively high during rim formation. We should expect that if WL rims grew by large increases in  $P_{\rm Mg}$  that there should be evidence for a comparable rise in  $f_{\rm O2}$ . The evidence for this increase in  $f_{\rm O2}$  is discussed next.

# 6. Oxidation state of Wark-Lovering rims

Titanium is usually present in terrestrial and extraterrestrial samples as Ti<sup>4+</sup>. In CAIs there is appreciable Ti<sup>3+</sup> as well as Ti<sup>4+</sup> in pyroxenes. The presence of

Table 2
Representative pyroxene compositions for CAI 144A

Analysis	WL rim		WL rim		Interior				
Measured	wt.% $\sigma_{\rm m}$		wt.% $\sigma_{\rm m}$		wt.% $\sigma_{\rm m}$				
SiO <sub>2</sub>	42.26	0.13	34.62	0.12	31.60	0.11			
$Al_2O_3$	16.71	0.08	23.54	0.09	19.69	0.09			
TiO <sub>2</sub>	2.50	0.05	7.66	0.08	17.85	0.13			
Ti <sub>2</sub> O <sub>3</sub>	0.00	0.00	0.00	0.00	0.00	0.00			
FeO	1.32	0.04	1.08	0.04	0.00	0.00			
Cr <sub>2</sub> O <sub>3</sub>	0.09	0.04	0.19	0.04	0.08	0.04			
MnO	0.00	0.00	0.00	0.00	0.00	0.03			
MgO	11.77	0.06	10.14	0.06	7.04	0.05			
CaO	24.17	0.10	23.54	0.10	24.52	0.10			
Na <sub>2</sub> O	0.11	0.02	0.05	0.02	0.00	0.00			
$K_2O$	0.04	0.02	0.04	0.02	0.01	0.01			
Total	98.96		100.86		100.80				
Cations formula based on 6 oxygen atoms									
Si	1.559		1.267		1.172				
Al	0.727		1.016		0.861				
Ti <sup>4+</sup> =Total	0.069		0.211		0.498				
Fe	0.041		0.033		0.000				
Cr	0.003		0.005		0.002				
Mn	0.000		0.000		0.000				
Mg	0.647		0.553		0.389				
Ca	0.955		0.923		0.974				
Na	0.008		0.004		0.000				
K	0.002		0.002		0.000				
Total	4.011		4.014		3.896				
Recalculated	wt.%	$\sigma_{\mathrm{m}}$	wt.%	$\sigma_{ m m}$	wt.%	$\sigma_{\mathrm{m}}$			
$SiO_2$	42.26	0.13	34.62	0.12	31.61	0.11			
$Al_2O_3$	16.71	0.08	23.55	0.09	19.70	0.08			
$TiO_2$	3.72	0.27	9.21	0.26	6.88	0.22			
$Ti_2O_3$	-1.10	0.25	-1.39	0.24	9.87	0.22			
FeO	1.33	0.04	1.08	0.04	0.00	0.00			
$Cr_2O_3$	0.09	0.04	0.19	0.04	0.08	0.04			
MnO	0.00	0.00	0.00	0.00	0.00	0.03			
MgO	11.77	0.06	10.14	0.05	7.04	0.05			
CaO	24.17	0.11	23.55	0.10	24.52	0.10			
Na <sub>2</sub> O	0.11	0.02	0.05	0.02	0.00	0.00			
K <sub>2</sub> O	0.04	0.02	0.04	0.02	0.01	0.01			
Total	99.09		101.03		99.71				
Tetrahedral ca Si	tions 1.555	0.004	1.263	0.004	1.203	0.004			
Al	0.445	0.004	0.737	0.004	0.797	0.004			
Total	2.000	0.003	2.000	0.003	2.000	0.003			
10141	2.000		2.000		2.000				
Octahedral cat Al		0.002	0.275	0.002	0.007	0.002			
Al Ti <sup>4+</sup>	0.280	0.003	0.275	0.003	0.087	0.003			
Ti <sup>3+</sup>	0.103	0.008	0.253	0.007	0.197	0.006			
	-0.034	0.008	-0.042	0.007	0.314	0.007			
Fe Cr	0.041	0.001	0.033	0.001	0.000	0.000			
Cr	0.003	0.001	0.005	0.001	0.002	0.001			

Table 2 (continued)

Analysis	WL rim		WL rin	ı	Interior	
Measured	wt.%	$\sigma_{ m m}$	wt.%	$\sigma_{ m m}$	wt.%	$\sigma_{ m m}$
Octahedral	cations					
Mn	0.000	0.000	0.000	0.000	0.000	0.001
Mg	0.645	0.003	0.551	0.003	0.399	0.002
Ca	0.953	0.003	0.920	0.003	1.000	0.003
Na	0.008	0.001	0.004	0.001	0.000	0.000
K	0.002	0.001	0.002	0.001	0.000	0.000
Total	2.001		2.001		1.999	
$Ti^{3+}/Ti^{4+}$	-0.33	0.05	-0.17	0.03	1.60	0.08
$\log f_{\mathrm{O2}}$ –	14.53	0.64 -	- 13.01	0.60 -	- 19.26	0.10

Measurements are representative WL rim and interior pyroxene compositions. The amount of  ${\rm Ti}^{3+}$  is calculated from  ${\rm Ti}^{4+}$  based on oxygen excess in the cation-normalized formula. Reported uncertainties are  $1\sigma_{\rm m}$  based on Monte Carlo error simulation (n=300 trials, see text for details).

 ${
m Ti}^{3+}$  signifies reducing conditions similar to solar values of  $f_{\rm O2}$  [8]. In this study  $f_{\rm O2}$  values for the rim and interior of CAI 144A were determined using electron microprobe analyses of Ti-bearing pyroxene grains (representative analyses can be seen in Table 2). The ratio of the activity of  ${
m CaTi}^{4+}{
m Al}_2{
m O}_6$  to the activity of  ${
m CaTi}^{3+}{
m AlSi}{
m O}_6$  in pyroxene defines  $f_{\rm O2}$  by virtue of gas—solid equilibria involving  ${
m O}_2$ . For pyroxene in the interior of the CAI where melilite—gas equilibria calibrated previously by Beckett and Grossman [8]. The first is the reaction

$$\begin{split} & 4 Ca Mg Si_{2}O_{6} \ + 4 Ca Ti^{4+}Al_{2}O_{6} \\ & = 4 Ca Ti^{3+}Al SiO_{6} \ + 2 Ca_{2} Mg Si_{2}O_{7} \\ & \quad + 2 Mg Al_{2}O_{4} \ + O_{2} \\ & \quad Sp \end{split} \tag{3}$$

and the second is

$$\begin{split} &2CaAl_{2}SiO_{6}\ + 2CaMgSi_{2}O_{6}\ + 4CaTi^{4+}Al_{2}O_{6}\\ &= 4CaTi^{3+}AlSiO_{6}\ + 2Ca_{2}Al_{2}SiO_{7}\\ &\quad + 2MgAl_{2}O_{4}\ + O_{2}\\ &_{Sp} \end{split} \tag{4}$$

In both reactions di, Cats, T4Cats and T3Cats refer to the indicated components in pyroxene and ak and gh to the components in melilite. In the WL rims melilite is scarce to absent and we employ a different reaction involving pyroxene and gas alone:

$$\begin{split} &2 Ca Mg Si_2 O_6 \ + 4 Ca Ti^{4+} Al_2 O_6 \ + 2 Si O \\ &di \\ &= 2 Ca Al_2 Si O_6 \ + 4 Ca Ti^{3+} Al Si O_6 \ + 2 Mg \ + O_2 \ . \\ &Cats \\ &T3 Cats \\ & &Cats \\ \end{split} \ . \label{eq:catalog} \tag{5}$$

A systematic study of the accuracy of the thermodynamic data for the Ti-bearing components in pyroxene has not been undertaken. The uncertainties in these thermochemical data are removed in the case of reaction (5) if the equilibrium constant is expanded in differential form. The equilibrium for reaction (5) leads to the expression

$$\log f_{O2} = 4\log\left(\frac{a_{T4Cats}}{a_{T3Cats}}\right) + 2\log\left(\frac{a_{di}}{a_{Cats}}\right) + 2\log\left(\frac{P_{SiO}}{P_{Mg}}\right) - \frac{\Delta G_{rxn}^{0}}{2.3RT}$$
(6)

where  $\Delta G_{\rm rxn}^0$  is the Gibbs free energy for the reaction at a standard state of pure solids and ideal gases at 1 bar, R is the ideal gas constant, T is temperature, and  $a_i$  is the activity of the indicated component i. The influence of the uncertainties in  $\Delta G_{\rm rxn}^{\ \ 0}$  is removed by taking the derivative of the  $\log f_{\rm O2}$  in (6), recognizing that the coefficients on the right-hand side are the partial derivatives of  $\log f_{\rm O2}$  with respect to the logarithms of the indicated activity ratios. The resulting expression is

$$\log f_{\rm O2} = \log f_{\rm O2}^0 + 4\Delta \log \left(\frac{a_{\rm T4Cats}}{a_{\rm T3Cats}}\right)$$

$$+ 2\Delta \log \left(\frac{a_{\rm di}}{a_{\rm Cats}}\right) + 2\Delta \log \left(\frac{P_{\rm SiO}}{P_{\rm Mg}}\right)$$
 (7)

where  $\log f_{O2}^{0}$  refers to some initial condition, in this case the  $\log f_{O2}$  of the CAI interiors. This last

equation permits contouring of log  $f_{\rm O2}$ -temperature space for  $\log(a_{\rm Ti4+}/a_{\rm Ti3+})$  in Ti-rich pyroxene in the absence of melilite (see Fig. 4). The problem is simplified further by recognizing that changes in  $P_{\rm Mg}$  will be attended by concomitant changes in  $P_{\rm SiO}$  (since the source of both is most likely silicate dust), meaning that the last term is insignificant where changes in orders of magnitude of  $f_{\rm O2}$  are of interest.

All three equilibria shown above were evaluated using activities based on ideal mixing on the octahedral site (in accord with the coupled mixing on the octahedral and tetrahedral sites in the pyroxenes and melilites associated with the tschermak exchange mechanism). In the cases of reactions (3) and (4) we used a fixed mole fraction of ak in melilite of 0.2 as indicated by electron microprobe analyses of melilites in the margin of Leoville CAI 144A. The results are robust relative to changes in ak content. Electron microprobe analyses were obtained from 22 Ti-rich pyroxene spots in the interior and from 29 relatively Ti-rich pyroxene spots in the WL rim of Leoville 144A. A complete data set is included in Appendix A, an electronic supplement.

Interior Ti-rich pyroxenes have an average Ti<sup>3+</sup> cations per 6 oxygens of  $0.31 \pm 0.01$   $2\sigma$  (Fig. 4), indicating  $f_{O2}$  values just below those defined by a solar gas [24]. In contrast, the Ti-rich pyroxenes of the WL rims yield a much lower average Ti3+ content of  $-0.03 \pm 0.02$  2 $\sigma$  cations per 6 oxygens with most points exhibiting no detectable Ti3+ (Slightly negative values for Ti3+ result from cation sums per 6 oxygens greater than 4. All such instances are within the uncertainties of the measurements). The IW  $f_{O2}$  buffer curve coincides with the loss of detectable Ti<sup>3+</sup> in pyroxene. As a result, absence of detectable Ti<sup>3+</sup> in the WL rims places a lower limit on the  $f_{O2}$  of their formation at or ~1 log units below the IW buffer. The lack of Ti3+ in the WL rims suggests that they formed at a minimum of ~6 to 7 log units higher  $f_{O2}$  than the CAI interiors (Fig. 4); the WL rims formed at  $f_{O2}$  indistinguishable from most chondrites, Earth, Mars, and Moon [25,26].

These results can be related back to the phase equilibrium model for WL formation in Fig. 3. The figure shows that WL rim formation could have been driven by a shift to higher  $P_{O2}$  ( $f_{O2}$ ) relative to the

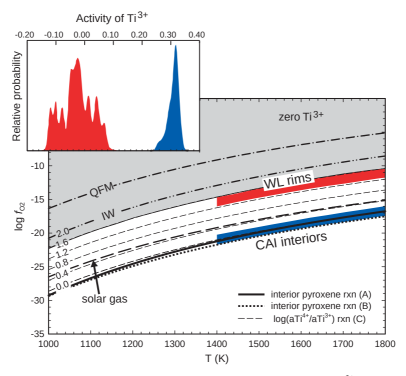


Fig. 4. Log  $f_{O2}$  versus temperature for average CAI interior and WL rim measurements based on  $Ti^{3+}$  in pyroxene. The inset shows the distribution of  $Ti^{3+}$  activity for interiors (near zero) and rims (~0.30). For substitution in the octahedral site of pyroxene the activity of  $Ti^{3+}$  is the mole fraction and therefore numerically equivalent to the number of  $Ti^{3+}$  atoms per 6 oxygen atoms (i.e., ideal mixing-on-sites activity model). Also shown are the Quartz–Magnetite–Fayalite (QFM) buffer, Iron–Wüstite (IW) buffer, and a curve representing a gas of solar composition. Calculated contours (light dashed lines) for the log of the activity ratio of T4Cats to T3Cats in 0.4 log unit intervals are shown for reference (see text for details).

CAI interiors. This requisite rise in  $f_{O2}$  is recorded by the pyroxenes in the rim of CAI 144A.

# 7. Rapid changing conditions in the solar nebula

Collectively, our observations imply a common nebular origin for WL rims by a process different from those that created the CAI interiors. Our findings provide evidence for two distinct episodes of nebular CAI evolution: (1) evaporation at low total pressure, low solar-like  $f_{\rm O2}$ , and low  $P_{\rm Mg}$  and (2) condensation to form the WL rims at high  $f_{\rm O2}$  (~IW) and high  $P_{\rm Mg}$ . The commonly accepted explanation for enhancing  $P_{\rm O2}$  and  $P_{\rm Mg}$  is evaporation of chondritic dust in the nebula [27]. The question arises as to whether these differences reflect changing conditions at a particular place in the nebula, as for example where CAIs remain in the hot inner nebula for long periods of

time (e.g., [28]) or transport from one environment to another (e.g., [29,30]). This work affords constraints on the applicability of these different models by comparing high-resolution age constraints with large shifts in ambient conditions during the formation of Wark–Lovering rims on CAIs.

The simple explanation for these data is that WL rims are condensates formed when CAIs passed from low pressures of a solar-like gas into regions of chondritic dust enrichment. Both the lack of Mg isotope fractionation and the high  $f_{\rm O2}$  are explained in this scenario. High  $P_{\rm O2}$  and  $P_{\rm Mg}$  are taken as evidence for dust enrichment relative to a solar gas. The initial  $^{26}{\rm Al}/^{27}{\rm Al}$  of  $5.3\times10^{-5}$  for the rims places temporal constraints on this movement from one reservoir to another if the  $(^{26}{\rm Al}/^{27}{\rm Al})_0$  for the solar system was uniform; the transition took 100,000 to 300,000 yr. Alternatively, these data constrain the degree of  $^{26}{\rm Al}$  heterogeneity in the solar nebula. CAIs and their rims

formed in markedly different conditions yet they have similar ( $^{26}\text{Al}/^{27}\text{Al}$ )<sub>0</sub> values of  $6\times10^{-5}$  to  $7\times10^{-5}$  and  $5.3\times10^{-5}$ , respectively. If the CAIs formed in the inner annulus of the protoplanetary disk near the *x*-point for example [31], where gas resembled a solar composition, and then were transported outwards to make WL rims in the part of the protoplanetary disk where chondrites and ultimately planets accreted, then the measured ( $^{26}\text{Al}/^{27}\text{Al}$ )<sub>0</sub> ratios limit the degree of heterogeneity in this parameter; the disparate regions of the nebula had a limited range in ( $^{26}\text{Al}/^{27}\text{Al}$ )<sub>0</sub> of  $\sim 7\times10^{-5}$  to  $\sim 5\times10^{-5}$ . In any case, the CAIs moved from a reducing solar-like gas to a more oxidizing region of chondritic dust enrichment in  $3\times10^5$  yr, or less.

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# Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.epsl. 2005.08.004.

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