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O, Mg, and Si isotope distributions in the complex ultrarefractory CAI Efremovka 101.1: Assimilation of ultrarefractory, FUN, and regular CAI precursors

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Abstract

Oxygen, magnesium, and silicon isotopic compositions in the mineralogically complex, ultrarefractory (UR) calciumaluminum-rich inclusion (CAI) E101.1 from the reduced CV3 chondrite Efremovka confirm that E101.1 is a compound CAI composed of several lithological units that were once individual CAIs, free-floating in the solar protoplanetary disk. Each precursor unit was found to have had its own thermal history prior to being captured and incorporated into the partially molten host CAI.

Four major lithological units can be distinguished on the basis of their isotopic compositions. (1) Al-diopside-rich sinuous fragments, hereafter sinuous pyroxene, are ¹⁶O-rich ($\Delta^{17}O \le -20\%$) and have light Mg and Si isotopic compositions with mass fractionation down to -3.5%/amu for both isotopic systems. We attribute these peculiar isotopic compositions to kinetic effects during condensation out of thermal equilibrium. (2) Spinel clusters are ¹⁶O-rich ($\Delta^{17}O \sim -22\%$) and have Mg isotope systematics consistent with extensive equilibration with the host melt. This includes (i) δ^{25} Mg values varying between + 2.6‰ and + 6.5‰ close to the typical value of host mellile at ~+5‰, and (ii) evidence for exchange of radiogenic ²⁶Mg with adjacent melilite as indicated by Al/Mg systematics. The spinel clusters may represent fine-grained spinel-rich proto-CAIs captured, partially melted, and recrystallized in the host melt. Al/Mg systematics indicate that both the sinuous pyroxene fragments and spinel clusters probably had canonical or near-canonical ²⁶Al contents before partial equilibration. (3) The main CAI host ($\Delta^{17}O \le -2\%$) had a complex thermal history partially obscured by subsequent capture and assimilation events. Its formation, referred to as the "cryptic" stage, could have resulted from the partial melting and crystallization of a ¹⁶O-rich precursor that underwent ¹⁶O-depletion and a massive evaporation event characteristic of F and FUN CAIs (Fractionated with Unknown Nuclear effects). Alternatively, a ¹⁶O-rich UR precursor may have coagulated with a ¹⁶O-poor FUN CAI having ⁴⁸Ca anomalies, as indicated by perovskite, before subsequent extensive melting. The Al/Mg systematics ($2.4 \times 10^{-5} \le (^{26}Al/^{27}Al)_0' \le 5.4 \times 10^{-5}$, where ($^{26}Al/^{27}Al_0'_0$ is a model initial ²⁶Al/²⁷Al ratio per analysis spot)

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are best understood if the FUN component was ²⁶Al-poor, as are many FUN CAIs. (4) A complete Wark-Lovering rim (WLR) surrounds E101.1. Its Mg and Si isotopic compositions indicate that it formed by interaction of the evaporated interior CAI with an unfractionated ¹⁶O-rich condensate component. Heterogeneities in ²⁶Al content in WLR spinels ($3.7 \times 10^{-5} \le ({}^{26}\text{Al}/{}^{27}\text{Al})'_0 \le 5.7 \times 10^{-5}$) suggest that the previously reported age difference of as much as 300,000 years between interior CAIs and their WLRs may be an artifact resulting from Mg isotopic perturbations, possibly by solid state diffusion or mixing between the interior and condensate components.

The isotopic systematics of E101.1 imply that ¹⁶O-rich and ¹⁶O-poor reservoirs co-existed in the earliest solar protoplanetary disk and that igneous CAIs experienced a ¹⁶O-depletion in an early high temperature stage. The coagulation of various lithological units in E101.1 and their partial assimilation supports models of CAI growth by competing fragmentation and coagulation in a partially molten state. Our results suggest that chemical and isotopic heterogeneities of unclear origin in regular CAIs may result from such a complex aggregation history masked by subsequent melting and recrystallization. © 2018 Elsevier Ltd. All rights reserved.

Keywords: Meteorites; CAIs; O isotopes; Mg isotopes; Si isotopes

1. INTRODUCTION

As the first solids to have formed in our solar system, Calcium-Aluminum-rich Inclusions (CAIs) from chondritic meteorites provide a snapshot of the state of the solar system at its very beginning, probably as it was a dense protoplanetary disk, when the protosolar cloud envelope was still collapsing (Yang and Ciesla, 2012; Mishra and Chaussidon, 2014; Taillifet et al., 2014). CAIs are of interest for astrophysics because their conditions of formation correspond to those prevailing in the region of telluric planet formation, within the first 1 AU from the protosun. However, in spite of almost five decades of intense study, the astrophysical context of CAI formation is still debated (e.g., Shu et al., 1997; Itoh and Yurimoto, 2003; Boss et al., 2012; Yang and Ciesla, 2012; Taillifet et al., 2014). Particular CAI properties that would be potentially useful for constraining astrophysical models include (1) their ¹⁶O and ²⁶Al isotopic anomalies (e.g., Itoh and Yurimoto, 2003; Mishra and Chaussidon, 2014; Aléon, 2016), (2) their high temperature gas-liquid-solid thermal histories (e.g., Richter et al., 2006; Shahar and Young, 2007), and (3) their redox state and the oxygen fugacity in the ambient medium where they formed (e.g., Simon et al., 2005, 2007; Grossman et al., 2008a; Paque et al., 2013). Up to now, most studies have been focused on one or two of these properties in isolation, however it has been recognized recently that many CAIs are compound objects that appear to have formed by aggregation of multiple precursors identified in the petrography (e.g., El Goresy et al., 2002; Aléon et al., 2007; MacPherson et al., 2012; Ivanova et al., 2015) or inferred from the isotopic record (e.g., Simon et al., 2017). Such CAIs most likely grew by coagulation-fragmentation either in the solid or in a partially molten state (Charnoz et al., 2015), thus adding complexity in the interpretation of differing petrological, chemical and isotopic properties in terms of astrophysical environments and processes that could produce mixing of refractory precursors.

The goal of our study is to use multiple tracers of the conditions of formation of CAIs on a well characterized compound object in order to disentangle the various steps of its formation history and to shed light on its precursors and on the astrophysical origin of the ¹⁶O and ²⁶Al isotopic

anomalies. We focused on Efremovka 101.1 (hereafter E101.1), a CAI previously studied in detail for its petrography, trace element and, to a limited extent, ²⁶Al content by El Goresy et al. (2002). Notably, it contains several lithological units interpreted to be CAI xenoliths trapped in a host inclusion. This petrographic stratigraphy provides an opportunity to unravel the sequence of events that led to the CAI formation, growth, and thermal processing, whereas in many large CAIs extensive melting has left only cryptic traces of original formation events. We performed an in-situ systematic O, Mg, Si isotope study of all lithological units coupled with complementary petrographic observations, in order to have five isotopic tracers (O, Mg and Si mass fractionations, ¹⁶O excesses and radiogenic ²⁶Mg excesses, hereafter ²⁶Mg*) to study the thermal history of all components, their relationships, and the interplay between thermal history and the record of the ¹⁶O and ²⁶Al isotopic anomalies.

2. SAMPLE AND METHODS

2.1. CAI Efremovka 101.1

E101.1 is a \sim 1.6 mm diameter Compact Type A (CTA) inclusion (Fig. 1a) from the Efremovka reduced CV3 chondrite, previously studied by El Goresy et al. (2002) for mineralogy, petrography, and Rare Earth Elements (REE) abundances. Details of its lithological units, including those defined by El Goresy et al. (2002) are given in supplementary information (Fig. S1) and listed in Table 1. Table 1 also summarizes the terminology and CAI-related acronyms used throughout the manuscript. E101.1 is dominated by melilite of typical composition between $Åk_{25}$ and $Åk_{30}$ and it contains numerous clusters of coarse-grained spinel (Fig. S1A). E101.1 exhibits two major properties that are uncommon among CAIs: (1) the presence, in the same half of the inclusion, of several lithological units suggesting incorporation of xenoliths that were once individual CAIs free-floating in the nebula (El Goresy et al., 2002, Fig. S1B), some of which are associated with oxidized secondary phases, and (2) systematic ultrarefractory (UR) REE patterns associated with presence of an aluminous clinopyroxene containing up to 12.9 wt% Sc₂O₃ and up to

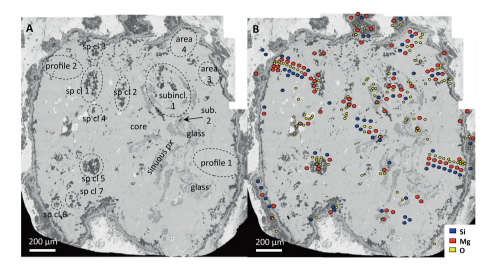


Fig. 1. Backscattered electron (BSE) images of CAI Efremovka 101.1. (a) Regions of interest. (b) Ion probe analysis spots.

Table 1
Abbreviations, types of pyroxene, lithological units and model endmembers discussed throughout the manuscript.

Denomination	Meaning										
Abbreviations											
AOA	Amoeboid Olivine Aggregate										
AR	Accretionary Rim (forsterite-rich in E101.1)										
WLR	Wark Lovering Rim										
CAI	Ca-Al-rich Inclusion										
UR	Ultra-Refractory										
FUN	Fractionated with Unknown Nuclear effects										
FG-sp-rich CAI	Fine-Grained spinel-rich CAI with nodular texture										
CMAS	$CaO + MgO + Al_2O_3 + SiO_2$ chemical system										
IMF	Instrumental Mass Fractionation										
RSF	Relative Sensitivity Factor										
Types of pyroxene											
WLR pyroxene	Ca-rich clinopyroxene with variable Al and Ti content ranging from very Al and Ti rich to pure diopside										
Sinuous pyroxene	Al-diopside										
Sc-Zr-rich pyroxene	Ca-rich clinopyroxene with elevated Al, Ti, Sc and Zr content and low SiO ₂ content										
Lithological units											
Sinuous pyroxene fragments	Sinuous fragments dominated by Al-diopside and locally containing FeO-rich assemblages										
Clusters 1–7	Coarse-grained spinel clusters in the main host										
Core	Central portion of the main host, mostly melilite										
Profiles 1 and 2	Chemical and O, Mg, Si isotopic traverses started from the rim and directed towards the center of the										
	inclusion, \sim perpendicular to the surface										
Areas 3 and 4	Regions between the sinuous pyroxene fragments and the rim, petrographically similar to those sampled by profile 1 and 2 but with different isotopic systematics										
Subinclusions 1 and 2	Subcircular melilite + spinel regions nearly enclosed in sinuous pyroxene, proposed to be xenoliths in El Goresy et al. (2002)										
Quenched glass	Regions with chemistry intermediate between melilite and pyroxene with Schlieren-like heterogeneities and dendritic melilite micro-crystals, located at the sinuous pyroxene - host melilite interface										
Model endmembers											
Step 1 parental melt	Ideal CMAS composition required to match the observed composition of intermediate areas (3, 4 and subinclusion 1) upon mixing with sinuous pyroxene										
Step 2 precapture melt	Ideal CMAS composition calculated from step 1 parental melt by substraction of condensate spinel										
Step 3 preevaporation composition	Ideal CMAS composition calculated from step 2 precapture melt by correction for evaporation based on Mg and Si isotopes										

 $5.4 \text{ wt}\% \text{ ZrO}_2$. E101.1 is surrounded by a complete Wark-Lovering rim (WLR, Wark and Lovering, 1977) and exhibits portions of a forsterite-rich accretionary rim (AR, Krot et al., 2001) in its outer part (Fig. S1C). The UR-REE patterns are found throughout the inclusion in all lithological units including both rims and the xenoliths. The most abundant xenoliths (more than 14) are made of sinuous fragments dominated by Al-diopside with chemistry similar to that often found in fine-grained spinel-rich CAIs or amoeboid olivine aggregates (Krot et al., 2004a, b) and locally including complex associations of finegrained FeO-rich minerals such as andradite and hedenbergite with wollastonite (El Goresy et al., 2002, Fig. S1B). Fine-grained symplectite intergrowths of spinel, clinopyroxene and perovskite, reminiscent of the reaction texture found in E49, another compound CAI from Efremovka (Aléon et al., 2007), are associated with the sinuous fragments and attributed to incorporation of small finegrained CAI fragments (El Goresy et al., 2002).

Two areas dominated by melilite and surrounded by sinuous pyroxene fragments have been defined as subinclusions (El Goresy et al., 2002, Fig. S1B). The host inclusion contains several populations of Y- and Zr-rich perovskite grains distinguished by their ZrO₂/Y₂O₃ ratios. The largest perovskite grains are commonly associated with FeNi metal. The Sc-Zr-rich pyroxene is systematically associated with perovskite and/or metal and defines reaction rims with the host melilite (El Goresy et al., 2002). Several regions at the interface between sinuous pyroxene and host melilite present melt pockets with compositions roughly intermediate between diopside and gehlenite (Fig. S1D). These melt pockets contain dendritic crystals of nearly pure gehlenite (Fig. S1D) and were described as an impact glass (El Goresy et al., 2002), hereafter referred to as the quenched glass. The innermost part of the host CAI is hereafter designated as the core. Three perovskites of differing ZrO_2/Y_2O_3 ratio were analyzed for Ca and Ti isotopes and show both excesses and depletions in ${}^{48}\text{Ca}$ (-7.5% < δ^{48} Ca < +9.8‰) but, remarkably, all have δ^{50} Ti $\approx 0‰$. This unusual property is also found in FUN-type (Fractionated with isotopic anomalies of Unknown Nuclear origin) and hibonite-rich CAIs from Murchison (Kööp et al., 2018). The three analyzed perovskite belong to three differing ZrO_2/Y_2O_3 populations and their $\delta^{48}Ca$ range is comparable to that of the FUN-type hibonites, apart from one having a δ^{48} Ca value of +43% (Kööp et al., 2018). El Goresy et al. (2002) reported early Mg isotopes measurements in spinel and melilite with a $\sim 1\%$ precision (1 σ) and in anorthite with a ~ 4 to $\sim 8\%$ precision (1 σ). Their Al/Mg systematics yielded a slightly sub-canonical isochron of slope ${}^{26}\text{Al}/{}^{27}\text{Al}_0 \sim 4.3 \times 10^{-5}$ in the sinuous fragments, scattered initial ²⁶Al/²⁷Al ratios in the host melilite and little to no ²⁶Mg excesses in anorthite, thus suggesting partial isotopic re-equilibration during or after ²⁶Al decay (El Goresy et al., 2002).

2.2. Petrographic analyses

Petrographic analyses were done using various electron probes and scanning electron microscopes (SEM) for comparison with isotopic analyses in the same phases. Maps of the distributions of Mg, Al, and Si in melilite were obtained using wavelength dispersive spectroscopy (WDS) with the Cameca SX5 and SX100 electron microprobes at the CAM-PARIS facility in the Pierre et Marie Curie University in Paris. Additional chemical analyses of melilite within 10 µm of oxygen isotope analyses pits were obtained with the SX100 electron probe of the CAMPARIS facility. The small extent of some gehlenitic regions, notably around spinel, compared to the spot size of O isotopes may lead to artificial discrepancies in the relationship between melilite chemistry and O isotopes in some cases. To minimize this effect, the Åk content was determined from the Åk map in the exact location of the SIMS spots and this Åk content was preferred over the spot analyses in these cases with strong chemical gradients (Fig. S2). All electron probe analyses are available as supplementary information. Backscattered electron (BSE) images of ion microprobe crater pits, were obtained using the SEMs at the University of California Los Angeles (UCLA) and in the Geosciences Paris Sud (GEOPS) laboratory at the Orsay University, and with the field emission gun SEM (FEG-SEM) in the Geology department at the Ecole Normale Superieure (ENS) in Paris.

2.3. Ion microprobe analysis of O, Mg, and Si isotopes

All isotopic analyses were conducted using the Cameca IMS 1270 ion microprobe at UCLA in two sessions for O isotopes, one session for Mg isotopes, and two sessions for Si isotopes (Fig. 1b). Issues related with instrumental biases such as differences in instrumental mass fractionation (IMF) due to chemical variations in the target (matrix effects) and analyses overlapping on different mineral phases are discussed in supplementary information. Uncertainties arising from these effects were taken into account and untrusted results were not considered in the discussion.

2.3.1. Oxygen isotopes

Oxygen isotope ratios were measured in multicollection mode using a $\sim 10 \,\mu m$ primary Cs⁺ beam of ~ 0.2 nA. Negative ions of ^{16}O , ^{17}O and ^{18}O were detected on an off-axis Faraday cup (FC), and on two electron multipliers (EMs, axial and an off-axis), respectively. The mass resolving power (MRP, defined as M/ Δ M) was set at ~6500, largely sufficient to resolve the ¹⁶OH⁻ interference at mass 17. The normal incidence electron gun was used for charge compensation. After a 30 s presputtering with a 0.5 nA beam, ¹⁶O, ¹⁷O and ¹⁸O were counted for 30 cycles of 10 s at typical count rates of 5×10^7 counts per second (cps), 2×10^4 cps and 1×10^5 cps for ¹⁶O, ¹⁷O and ¹⁸O respectively. Liquid nitrogen was used to keep the vacuum at $\sim 1 \times 10^{-9}$ Torr as a routine precaution but the plexiglas mount of E101.1 ensured limited outgassing. Data were corrected for IMF using a San Carlos olivine and a Burma spinel standard, which yielded similar IMF. All analyses were thus corrected with an average IMF. As a whole, matrix effects for O isotopes in the phases investigated here are negligible except for the silicate melt inclusion in metal and possibly for the FeO-rich silicates and the Sc-Zr-rich pyroxene (supplementary information). Results are reported using the standard δ -notation, where $\delta^{17,18}O =$ $(^{17,18}O/^{16}O_{samp} - ^{17,18}O/^{16}O_{SMOW})/(^{17,18}O/^{16}O_{SMOW}) \times$ 1000 with SMOW being the Standard Mean Ocean Water. Average 2σ uncertainties taking into account internal precision and reproducibility on standards are 0.48% and 0.94‰ on δ^{18} O and δ^{17} O, respectively. Deviations from the slope 0.52 terrestrial mass fractionation line are given as Δ^{17} O where Δ^{17} O = δ^{17} O – 0.52 × δ^{18} O. Mass fractionations are noted Δ^{18} O_{CCAM} and are calculated respectively as isotopic deviations from the classical slope 0.94 Carbonaceous Chondrite Anhydrous Minerals mixing line (Clayton and Mayeda, 1984) along a slope 0.52 line. They are given in ‰/atomic mass unit (‰/amu).

We obtained a total of 128 analyses, sampling most phases (melilite, spinel, forsterite, Al-diopside, FeO-rich assemblages, Sc-Zr-rich pyroxene, perovskite, impact glass, melt inclusion in metal, nepheline \pm anorthite) in several lithological contexts including the host CAI, sinuous pyroxene fragments, subinclusions 1 and 2, WLR and AR. Two profiles starting from the rim were done toward the interior of the host CAI (profiles 1 and 2) (Fig. S1E). The core of the host CAI was also targeted, as well as two areas between the sinuous pyroxene fragments and the rim (areas 3 and 4) (Fig. S1E). Both spinel and melilite inside of and in proximity to spinel clusters (clusters 1–5) were measured (Fig. S1A); in these regions melilite is significantly more gehlenitic than the typical host melilite.

2.3.2. Magnesium isotopes

Magnesium isotopes were measured using a \sim 30 µm O⁻ primary ion beam of 12–15 nA: ²⁷Al and the three Mg isotopes (²⁴Mg, ²⁵Mg and ²⁶Mg) were detected as positive ions in multicollection mode by using four Faraday cups. The MRP was set at ~4200, sufficient to completely separate molecular and doubly charged ion interferences (⁴⁸Ca²⁺ and ⁴⁸Ti²⁺). ²⁷Al⁺, ²⁶Mg⁺, ²⁵Mg⁺ and ²⁴Mg⁺ were counted for 20 cycles of 10 s after a 30–50 s presputtering for spinel, pyroxene and olivine and for 100 cycles for melilite analyses except for analyses 60, 61 and 71–78 (20 cycles).

Standards used for determination of the instrumental mass fractionation and elemental relative sensitivity factors were Burma spinel, a synthetic pyroxene glass with no ²⁶Mg excess (P0), olivine and pyroxene from San Carlos, and Madagascar hibonite. Measurements of standards were interspersed with sample analyses, thus defining 7 groups of data. For each group, standards data expressed as $1000 \times ln[({}^{25,26}Mg/{}^{24}Mg)_{measured})/({}^{25,26}Mg/{}^{24}Mg)_{true}] \text{ were }$ used to calculate an instrumental mass fractionation law, which was used to determine the ²⁶Mg excesses (noted $\delta^{26}Mg^*$) by difference with the expected ${}^{26}Mg$ due to mass fractionation only. This instrumental mass fractionation law was preferred to that resulting from CAI evaporation in nature (Davis et al., 2015) because the instrumental effects between the standards were found to be larger than the natural effects observed in E101.1, thus yielding a potentially larger source of uncertainty if improperly corrected. The inaccuracy resulting from this assumption is at most about 0.25% and is typically below 0.1% (Fig. S3). All isotopic results were subsequently reported in δ-notation as per mil deviations relative to the DSM3 terrestrial reference (Galy et al., 2003). Measurements as an unknown of a synthetic pyroxene glass having ${\sim}1\%~^{26}Mg$ excess (P1) yielded an average $^{26}\mathrm{Mg}$ excess of 1.071 \pm 0.080 % (2 σ). Al/Mg ratios were determined from the $^{27}\text{Al}^+/^{24}\text{Mg}^+$ ratios corrected using the relative sensitivity

factors (RSF) determined from Burma spinel (spinel), the P0 synthetic Al-Ti-pyroxene glass (Al-diopside, Sc-Zr-rich pyroxene, impact glass) and Madagascar hibonite (melilite). IMF for the ${}^{25}Mg/{}^{24}Mg$ ratios were calculated using Burma spinel for spinel analyses, the pyroxene P0 glass (or in one case the San Carlos pyroxene normalized to P0) for Aldiopside, Sc-Zr-rich pyroxene, impact glass and melilite and San Carlos olivine for forsterite. Burma spinel and P0 glass were previously measured relative to USNM forsterite, which composition relative to DSM3 was determined by Galy et al. (2003). We further assumed that San Carlos olivine has the same Mg isotopic composition as USNM forsterite. Given the Earth mantle origin of both olivines, the uncertainty resulting from this assumption is much smaller than our analytical precision. In the absence of a proper melilite standard for Mg isotopes we initially assumed that hibonite could be used as a standard for the Al/Mg ratio of melilite, a reasonable approximation within uncertainties, and that the P0 glass could be used to determine the Mg isotopes IMF for melilite. Further matrix effect investigations indicate a $5 \pm 0.5\%$ matrix effect between P0 and Al-rich melilite for Mg isotopes, which was retroactively applied to the sample melilite data. The additional uncertainty due to this correction has been propagated and is included in the reported dataset. Al/Mg ratios for forsterite were not corrected for RSF but were close enough to 0 (average $0.015\pm0.015,\,2\sigma)$ to have negligible effects on isochron regressions. Still, we note that the $Al^+/$ Mg⁺ ratios on the San Carlos olivine were significantly lower than this (average $2.88 \pm 0.06 \times 10^{-4}$, 2σ), suggesting that minor amounts of Al-rich material (e.g., Al-diopside) were incorporated in the analyses in agreement with SEM imaging of the AR. The precisions obtained for Mg isotopic ratios depend on the Mg content: 2σ errors for δ^{25} Mg incorporating both internal (within spot) and external reproducibility on standards range between 1.06 ‰ and 1.75 % for melilite and glass, and between 0.37 % and 0.89 % for other minerals. 2σ errors on $\delta^{26}Mg^*$ range between 0.06 and 0.95 ‰, again with larger uncertainties for the Al-rich, Mg-poor phases.

We obtained a total of 85 measurements of Mg isotope ratios from spinel, Al-diopside, Sc-Zr-rich pyroxene, impact glass and melilite. Care was taken to sample the host inclusion, the sinuous pyroxene fragments, subinclusion 1, the WLR and AR as well as five different spinel clusters. Adjacent melilite was measured in 4 of the 5 spinel clusters. Whenever possible, Mg isotopes spots were located within 20 μ m of the O isotope analysis pits for a direct comparison. For example, the Mg isotope profiles 1 and 2 were started from the rim and were made in parallel with the O isotope traverse (Fig. S1E).

2.3.3. Silicon isotopes

Silicon isotopes were measured in multicollection mode as negative ions with a $\sim 20 \ \mu m \ Cs^+$ primary ion beam of 1–1.7 nA. ²⁸Si⁻ and ³⁰Si⁻ were detected simultaneously on two Faraday cups. ²⁷Al⁻ was measured as well as a check on the composition of the analyzed mineral. The mass resolving power was set at 2400, sufficient to separate the ²⁹SiH⁻ interference at mass 30. The normal incidence

electron gun was used for charge compensation. Typical secondary ion intensities for ²⁸Si⁻ ranged between 4×10^7 cps in melilite to 7×10^7 cps in olivine. Si isotopic compositions are reported as δ^{30} Si per mil deviations relative to the NBS 28 international reference. With such conditions, a Poisson counting statistic $\leq \pm 0.1\%$ is obtained after a few minutes integration (30 cycles of 5 s acquisition time and 30 s of presputtering). The IMF was corrected by using San Carlos olivine, P0 pyroxene glass, San Carlos pyroxene and new melilite standards specifically prepared for Si isotope analyses (Marin-Carbonne et al., 2012). Matrix effects are particularly large for silicon isotopes in melilite (Knight et al., 2009), which ionize poorly and have large IMF. This matrix effect is dependent on the Al content and was calibrated using the three melilite standards (Åk₁₀, Åk₈₀ and a zoned crystal with Åk content ranging between 15 and 70 mol%) and the method described in Marin-Carbonne et al. (2012). Possible matrix effects in pyroxene remained within analytical uncertainty (supplementary information). Typical errors after all corrections including propagation of internal precision and external reproducibility determined on standards are 0.4–0.6 % (2 σ).

The mass fractionation in %c/amu is expressed as FSi, where $FSi = \delta^{30}Si/0.501$ where the factor 0.501 is the (M²⁹Si-M²⁸Si)/(M³⁰Si-M²⁸Si) ratio, where M^xSi is the exact mass of the ^xSi isotope. Knight et al. (2009) report a mass fractionation factor of 0.518 ± 0.006 between δ^{29} Si and δ^{30} Si in evaporated CMAS (Ca, Mg, Al, Si) glasses analogs of CAIs, so that FSi as defined here is not strictly identical to δ^{29} Si, however the difference is small with regard to our uncertainties so that it does not affect any interpretation and FSi can be compared with δ^{25} Mg. Si isotopic analyses were acquired for 60 spots, preferentially in the vicinity of O and Mg isotope crater pits in order to allow as much intercomparison between the various isotopic systems as possible (within 20-30 µm). For example, Si isotope traverses were made in parallel with the O and Mg isotope profiles (Fig. S1E). Analyzed phases include olivine, pyroxene and melilite from different petrographic contexts of E101.1, including the host inclusion, sinuous pyroxene fragments, subinclusion 1, areas 3 and 4, and the WLR and AR.

3. RESULTS

3.1. Petrographic observations

New electron probe analyses indicate that in spite of a typical melilite composition in the Åk₂₀-Åk₃₀ range (El Goresy et al., 2002), melilite can reach nearly pure gehlenite composition (down to $Åk_3$) in contact with spinel, notably near the WLR as in other CAIs (e.g., Fahey et al., 1987; Goswami et al., 1994; Toppani et al., 2006; Aléon et al., 2007; Kawasaki et al., 2012), but around spinel clusters in the interior as well (Fig. S4). These interior spinel clusters correspond to the end of spinel trails reaching the rim of the inclusion. The electron probe mapping of Mg further shows that these spinel trails are systematically associated with aluminous melilite thus defining continuous to semicontinuous channels of aluminous melilite between the spinel clusters and the edge of the CAI (Fig. 2). This is notably the case of spinel clusters 1, 2 and 4 (Fig. 2). Being closer to the rim, cluster 3 is embedded in such a channel. Spinel clusters in the upper part of subinclusion 1 are also located at the end of such trails (Figs. 1 and 2). Cluster 5 and the smaller clusters in the lower left part of Fig. 1 also define a swarm of spinel associated with Al-rich melilite and connected to the edge of the CAI, although not as continuous as the upper trails (Fig. 2). Two of these small clusters are labeled as clusters 6 and 7 (Fig. 3). Detailed SEM examination of the spinels throughout the inclusion and especially of those connected to the rim via gehlenitic melilite reveals numerous signs of spinel resorption, which notably include (1) rounded anhedral crystal shapes (Fig. 3a and b), (2) embayments (Fig. 3a-c), (3) a rim around spinel consisting of Al-, Ti-clinopyroxene \pm spinel symplectite, locally with perovskite inclusions (Fig. 3). A mineralogical continuum is observed in these rims from pyroxene + abundant microspinel to pyroxene only to pyroxene + perovskite. The Ti content of the pyroxene can be used to map the symplectite (Fig. 3d). This symplectite is similar to that previously described in E49, another compound CTA inclusion from Efremovka (Aléon et al., 2007). Where the surrounding rim consists of clinopyroxene only, it is strongly reminiscent

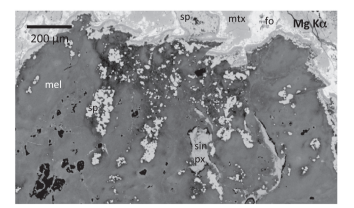


Fig. 2. Mg K α X-ray map of the upper portion of E101.1 in log scale. Spinel clusters are connected to the rim of the CAI by trails of spinel grains surrounded by gehlenitic melilite (dark). Black grains are devoid of Mg (perovskite, metal, anorthite, nepheline...). Abbreviations: sp - spinel, sin px - sinuous pyroxene, mel - melilite, fo - forsterite, mtx - matrix.

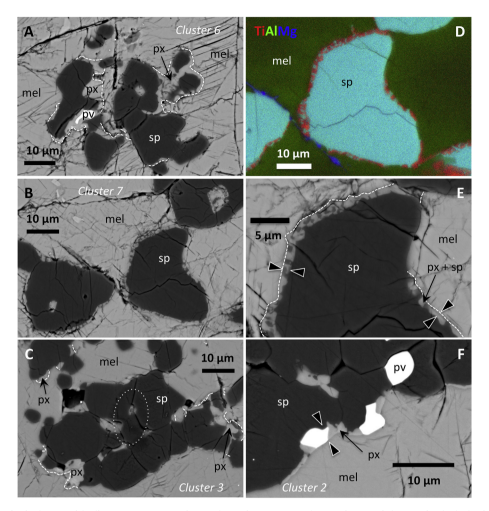


Fig. 3. Example of spinels (sp) with clinopyroxene (cpx) rims and reaction textures. (A) BSE image of cluster of anhedral spinel, embayed and rimmed by cpx. (B) BSE image of anhedral spinel grains almost completely enclosed in the cpx + sp rim symplectite detailed in D and E. (C) Cluster 3 dominated by subhedral spinels also contains grains with local embayments, anhedral shapes, and cpx \pm sp rim. BSE image, dotted ellipse indicates ion probe spot for Mg isotope analysis Mg3. (D) False color RGB TiAlMg map illustrating the distribution of the Ti-bearing cpx rim (red) with spinel µm and sub-µm inclusions around coarser spinel (light blue) in (B). (E) Detail of the cpx + sp rim around spinel in (B) and (D), BSE image. (F) Detail of the perovskite-containing cpx rim around spinel from cluster 2. Dashed white lines in (A), (C) and (E) and arrows in (E) and (F) indicate the border of the cpx \pm sp rim. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

of the boundary clinopyroxene described around spinel in Type B CAIs and in the mantles of Type B1 CAIs (Paque et al., 2009). This rim is also similar in textural relationship to the Sc-Zr-rich pyroxene rims around perovskite and metal grains (Fig. 4, El Goresy et al., 2002). As in the case of spinels, the perovskite rimmed by the Sc-Zr-rich pyroxene commonly shows embayments and anhedral shapes. FeNi metal is commonly associated with perovskite and rimmed by the Sc-Zr-rich pyroxene. It is predominantly high Ni taenite (35-51 at% Ni) but high Co kamacite (up to 8 at% Co) is also found, usually in heterogeneous grains (see supplementary informations). One grain contains 5-6 at% platinum group elements. Most grains contain inclusions of Fe-V-oxides and Ca-phosphates near their periphery. A troilite inclusion is found once. Three Si-rich glass inclusions were found in one of the metal grains (Fig. 4a).

Although E101.1 was initially described as mostly unaltered, we found nepheline associated with anorthite in replacement of melilite, notably in the vicinity of perovskite + metal nodules (Fig. 4).

3.2. Oxygen isotopes

Oxygen isotope ratios in E101.1 (Table S1) plot along the Carbonaceous Chondrites Anhydrous Minerals mixing line (Clayton and Mayeda, 1984) and span the whole range between the typical ¹⁶O-rich composition of spinel in CAIs ($\Delta^{17}O \sim -22\%$) and a sub-terrestrial ¹⁶O-poor composition (up to $\Delta^{17}O \sim -2\%$, Fig. 5a).

Spinel and accretionary rim forsterite are uniformly ¹⁶Orich with $\Delta^{17}O \leq -20.5\%$. The isotopic composition of melilite varies from significantly enriched in ¹⁶O ($\Delta^{17}O$ down to -19.8% in the WLR, Fig. 5c) to relatively ¹⁶Opoor ($\Delta^{17}O$ up to -1.9‰). The most ¹⁶O-rich melilite is systematically gehlenitic melilite associated with spinel (Fig. 6), although the reverse is not systematic. This is true for

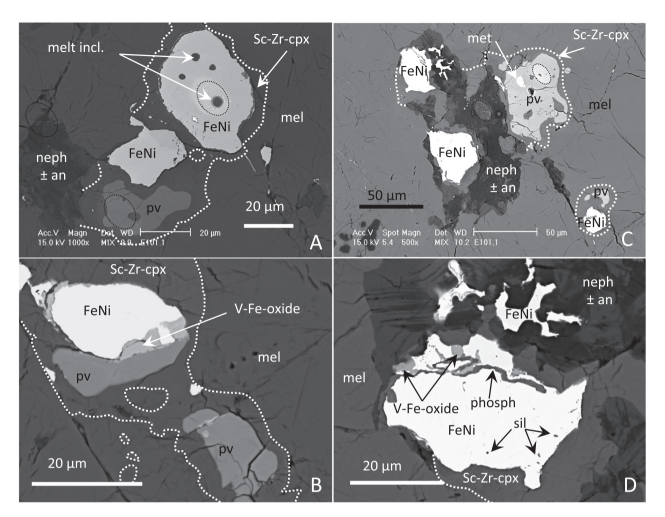


Fig. 4. BSE images of perovskite-metal (pv-met) nodules rimmed by the Sc-Zr-rich pyroxene. (A) nodule with silicate melt inclusions within metal. (B) V-Ti oxide in pv-met association. (C) Perovskite Per B. (D) Detail of top metal grain in C, showing metamorphic exsolution of silica, phosphate and V-Fe-oxide. Sc-Zr-rich pyroxene shown with dotted lines. Dotted ellipses indicate O isotope SIMS spots. Abbreviations: met - metal, pv - perovskite, cpx - pyroxene, mel - melilite, neph - nepheline, an - anorthite, phosph - phosphate, sil - silica.

melilite from the WLR and also for gehlenitic melilite associated with coarse-grained spinel clusters in the interior (Fig. 6b). Interestingly, the Ak content of melilite associated with spinel clusters define a trend similar to that in melilite as a function of distance to WLR spinel in profiles 1 and 2 (Fig. 6c). Examination of melilite composition in the various petrographic areas indicates that different regions in the interior of the CAI have statistically different O isotopic compositions (weighted mean ranging from $\Delta^{17}O = -5.01 \pm 0.27\%$ to $-8.64 \pm 0.43\%$, 2σ , Table 2). The most ¹⁶O-enriched regions are encountered in subinclusion 1 and areas 3 and 4, all in the vicinity of the sinuous pyroxene fragments. The reduced χ^2 for Δ^{17} O within each region ranges from 1.8 to 4.7. By contrast, the average $\Delta^{17}O$ value over all standard analyses is -0.18 ± 0.16 (2σ) with a reduced χ^2 of 1.9. This suggests that melilite in individual regions is not fully homogenized (Table 2). A small amount of mass fractionation relative to the CCAM line (weighted mean $\Delta^{18}O_{CCAM} = 2.33 \pm 0.16\%$ / amu, 2σ) is observed in the ¹⁶O-poor melilite but not in

the ¹⁶O-rich melilite (weighted mean $\Delta^{18}O_{CCAM} = 0.76 \pm 0.57\%$ /amu, 2 σ , Fig. 7). The amplitude of this effect suggests it is not of instrumental origin (supplementary information). The most ¹⁶O-depleted compositions ($-2.8\% \leq \Delta^{17}O \leq -1.9\%$) were observed in melilite in the outer portion of profile 1 (see below) and also in (1) nepheline, (2) Sc-Zr-rich pyroxene and (3) one glass inclusion in metal (Fig. 5d).

The Sc-Zr-rich pyroxene $(-3.26\% \le \Delta^{17}O \le -2.23\%)$ appears to be mass fractionated similarly to the ¹⁶O-poor melilite (weighted mean $\Delta^{18}O_{CCAM} = 2.49 \pm 0.75\%$ /amu, 2σ , Fig. 7) but the ¹⁶O-rich pyroxenes (Al-rich diopside from the sinuous fragments and pyroxene from the WLR) are not appreciably mass fractionated (weighted mean $\Delta^{18}O_{CCAM} = 0.64 \pm 0.29\%$ /amu, 2σ , Fig. 7). Again, this suggests that the apparent mass fractionation is not of instrumental origin although a matrix effect between diopside and Sc-Zr-rich pyroxene cannot be excluded (supplementary information). Many analyses of the Al-diopside in the sinuous fragments overlap with FeO-rich silicates

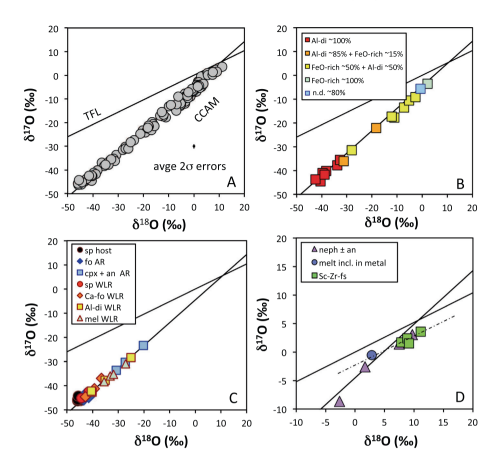


Fig. 5. O isotopic compositions in E101.1. (A) All data. (B) Sinuous pyroxene and associated FeO-rich silicates. Shades of color indicate different degree of mixing between Al-diopside and FeO-rich minerals within SIMS spots (proportions given). n.d. stands for not determined, most likely anorthite or possibly wollastonite. (C) minerals in the accretionary rim (AR) and Wark-Lovering rim (WLR). (D) relatively ¹⁶O-poor minerals in E101.1; dash-dotted line is a mass fractionation line through the most ¹⁶O-poor data. TFL - terrestrial mass fractionation line, CCAM - Carbonaceous Chondrites Anhydrous Minerals mixing line.

and show intermediate ¹⁶O-excess (Fig. 5b) between pure Al-diopside ($\Delta^{17}O \leq -20\%$) and FeO-rich silicates ($\Delta^{17}O \geq -5\%$). Pure Al-diopside analyses show a correlation between diopside chemistry and ¹⁶O-enrichment, with the most refractory pyroxene (with highest Al₂O₃ and lowest SiO₂ contents) being also the most ¹⁶O-rich (Fig. 8).

Perovskite was found to be variably enriched in ¹⁶O $(-19\% \le \Delta^{17}O \le -9\%)$ but the ¹⁶O-excess does not appear to be related to the ZrO₂/Y₂O₃ ratio, which was used by El Goresy et al. (2002) to characterize several populations of perovskite (Fig. 9).

Minerals in the WLR and the forsterite-rich accretionary rim are rich in ¹⁶O as often observed in CAIs (Krot et al., 2002; Yoshitake et al., 2005; Aléon et al., 2007; Bodénan et al., 2014). Apart from one analysis of WLR pyroxene ($\Delta^{17}O = -15.3 \pm 0.7\%$), a few analyses with intermediate composition ($-18\% \le \Delta^{17}O \le -13\%$) correspond to WLR melilite and pyroxene + anorthite nodules in the AR.

Finally the quenched glass appears to have oxygen isotopic compositions between those of the host melilite and Al-diopside from the sinuous pyroxene fragments (-14% $\leq \Delta^{17}O \leq -10\%$).

3.3. Magnesium isotopes

3.3.1. Initial ²⁶All²⁷Al ratios

A trend of increasing δ^{26} Mg^{*} with ²⁷Al/²⁴Mg is observed but a single well-defined isochron cannot be identified (Fig. 10, Table S2). A spread in δ^{26} Mg^{*} beyond analytical uncertainties is observed in melilite and spinel (Fig. 10). In order to evaluate the spread in excess ²⁶Mg and its consequence on the ²⁶Al distribution in the inclusion and on chronology, model initial ²⁶Al/²⁷Al, denoted (²⁶Al/²⁷Al)'₀ ratios have been calculated for individual data points. The (²⁶Al/²⁷Al)'₀ ratio is defined as the slope of an isochron going through a given analysis and through the origin.

In the interior, spinels from subinclusion 1, cluster 1 and cluster 5 have $\binom{2^{6}\text{Al}/^{27}\text{Al}}_{0}$ ratios consistent with a canonical ratio (i.e., 5.23×10^{-5} , Jacobsen et al., 2008) within error (weighted means of $(5.0 \pm 0.4) \times 10^{-5}$, $(5.3 \pm 0.9) \times 10^{-5}$, $(5.6 \pm 0.4) \times 10^{-5}$, 2σ , respectively, Table 2), but cluster 2 has a distinctly supracanonical weighted mean $((7.3 \pm 0.9) \times 10^{-5}, 2\sigma, \text{Table 2})$. Near the CAI periphery, cluster 3 has a subcanonical $\binom{2^{6}\text{Al}/2^{7}\text{Al}}_{0}$ ratio of $(3.6 \pm 1.2) \times 10^{-5}$ (2σ , Table 2).

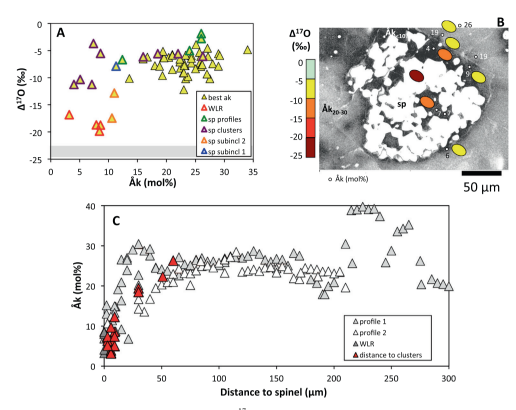


Fig. 6. Oxygen isotopic compositions of melilite in E101.1. (a) Δ^{17} O as a function of melilite chemistry. Grey box: typical value of spinel. 2σ errors. (b) Chemical map of melilite computed from Mg and Al X-ray maps. Bright: akermanitic mel, dark: gehlenite, spinel overlaid in white. Ellipses are ion probe spots with color coded Δ^{17} O value. (c) Chemistry of melilite as a function of distance to spinel. Profile 1, 2 and WLR are reported relative to WLR. Melilite in the vicinity of spinel clusters is reported relative to spinel in the corresponding cluster. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

Individual (²⁶Al/²⁷Al)₀ ratios in melilite range between $(5.4 \pm 0.2) \times 10^{-5}$ and $(2.4 \pm 0.5) \times 10^{-5} (2\sigma)$ suggesting secondary perturbations owing to Mg diffusion and isotope exchange in an initially canonical melilite. No clear difference can be observed between different regions in the CAI (Fig. 10), with similar average and spread in $({}^{26}\text{Al}/{}^{27}\text{Al})_0$ ratios of 3.7 ± 0.5 , 4.7 ± 0.6 , 4.1 ± 0.9 , 4.2 ± 0.3 , $4.0 \pm$ 0.5, 3.4 ± 0.7 , 4.0 ± 0.7 for area 3, area 4, subinclusion 1, core, melilite in the vicinity of spinel clusters, profile 1 and profile 2, respectively (weighted mean given in units of 10^{-5} ; uncertainties are 1 standard deviation here rather than 2 standard errors to take into account heterogeneities in melilite within individual areas and to emphasize the similarity of the ratios). In subinclusion 1 and spinel clusters 1 and 5, melilite associated with spinel have comparable $({}^{26}\text{Al}/{}^{27}\text{Al})'_0$ ratios between 4.1×10^{-5} and 4.3×10^{-5} , while associated spinels have higher ratios consistent with the canonical ratio (Table 2). Only melilite associated with sub-canonical spinel (cluster 3) has a (²⁶Al/²⁷Al)₀' ratio comparable to that of the associated spinel ((3.3 \pm 0.3) \times 10^{-5} and $(3.6 \pm 1.2) \times 10^{-5}$, respectively, 2σ , Table 2). This corresponds to systematic sub-canonical slopes and elevated intercepts in Al/Mg isochrons, with the highest intercept for cluster 2 (Fig. 11), except for cluster 3 where the intercept is consistent with 0. Regressions for subinclusion 1, cluster 1 and cluster 5 are identical within error (Fig. S5).

Due to large uncertainties, all estimates of the initial ${}^{26}\text{Al}/{}^{27}\text{Al}$ ratio of the sinuous pyroxene are consistent with the canonical ratio (Fig. 10) and with the average melilite value at the 2σ level.

In the WLR, $({}^{26}\text{Al}/{}^{27}\text{Al})_0'$ ratios range from canonical $((5.7 \pm 0.4) \times 10^{-5})$ to distinctly subcanonical $((3.7 \pm 0.5) \times 10^{-5})$ in spinel. If both spinel and pyroxene are considered together a regression yields a slope of $(4.5 \pm 0.8) \times 10^{-5}$ (2σ), which becomes $(4.8 \pm 0.5) \times 10^{-5}$ (2σ) if forced through the origin. This is identical to the weighted mean of $({}^{26}\text{Al}/{}^{27}\text{Al})_0'$ ratios of $(4.8 \pm 0.2) \times 10^{-5}$ (2σ , Table 2). Given that variations in spinel should affect the determination of the slope, we also investigated the ${}^{26}\text{Al}/{}^{27}\text{Al}$ systematics of the WLR without spinel. Pyroxene data are individually consistent with the canonical ratio and a regression yields a slope of $(4.1 \pm 3.0) \times 10^{-5}$ (2σ), which becomes $(6.0 \pm 0.9) \times 10^{-5}$ (2σ) if forced through the origin.

The weighted mean of ²⁶Mg excesses ($\delta^{26}Mg^*$) in accretionary rim forsterite is $0.06 \pm 0.04\%$ (2 σ), which indicates a small but significant excess of ²⁶Mg. The high Al⁺/Mg⁺ ratios compared to San Carlos olivine suggest Al-rich contamination in the analysis spot, so that this excess is probably intermediate between that of pure olivine and that of the bulk AR. It compares well with previous values in bulk AOA and Fo-rich AR in CAIs (Larsen et al., 2011) as well

	$\Delta^{17}O(\%)$					δ ²⁵ Mg (‰)					$({}^{26}\text{Al}/{}^{27}\text{Al})_0' [10^{-5}]^{**}$					FSi (‰)				
	#	min	max	wm	wm 2σ	#	min	max	wm	wm 2σ	#	min	max	wm	wm 2σ	#	min	max	wm	wm 2σ
Profile 1 Mel	13	-7.44	-1.86	-5.01	0.27	9	3.69	6.28	5.04	0.57	9	2.44	4.52	3.43	0.13	8	4.98	6.73	6.29	0.15
Profile 2 Mel	10	-9.32	-5.73	-6.93	0.31	10	1.88	6.91	5.06	0.55	10	2.91	4.97	3.95	0.11	9	5.16	6.56	5.66	0.13
Core Mel	3	-6.6	-5.12	-6.07	0.57	2	3.99	5.01	4.50	1.23	2	3.98	4.36	4.15	0.48	4	5.19	7.66	6.59	0.20
Area 3 Mel	5	-10.18	-7.28	-8.64	0.43	5	0.88	2.01	1.43	0.47	5	2.83	4.09	3.70	0.11	1			1.24	
Area 4 Mel	5	-9.59	-7.42	-8.03	0.4	5	0.54	2.15	1.49	0.47	5	3.91	5.39	4.71	0.11	3	2.8	3.53	3.25	0.25
Sub 2 Mel	2	-17.45	-12.82	-15.14																
Sub 1 Sp						3	-0.77	0.58	-0.02	0.48	3	4.53	5.84	5.04	0.37					
Sub 1 Mel	6	-12.18	-7.8	-8.78	0.36	4	-1.51	1.06	-0.12	0.53	4	2.96	5.04	4.07	0.13	3	3.99	5.69	4.99	0.26
Cluster 1 Sp	1			-22.83		3	6.42	6.70	6.53	0.51	3	4.93	5.72	5.30	0.89					
Cluster 1 Mel [*]	2	-10.31	-3.68	-7.15	0.71	2	5.18	5.71	5.47	0.78	2	4.00	4.53	4.22	0.17					
Cluster 2 Sp	1			-22.46		2	4.81	5.29	5.05	0.62	2	6.46	7.56	7.30	0.90					
Cluster 2 Mel*	1			-6.14		2	5.02	5.77	5.40	0.75	2	3.62	3.90	3.77	0.18					
Cluster 3 Sp						1			4.28		1			3.62						
Cluster 3 Mel*						1			6.10		1			3.34						
Cluster 4 Sp						1			6.63		1			5.30						
Cluster 4 Mel [*]																				
Cluster 5 Sp	1			-21.77		2	2.61	2.65	2.63	0.52	2	5.47	5.74	5.62	0.35					
Cluster 5 Mel*	6	-11.25	-5.51	-7.33	0.43	2	3.65	6.09	4.87	0.75	2	4.07	4.57	4.32	0.20	2	5.6	5.83	5.71	0.31
Sin Px	7	-23.56	-18.53	-20.98	0.36	4	-4.65	-2.39	-3.40	0.31	4	4.42	13.20	5.21	1.28	3	-4.48	-2.37	-3.09	0.37
Sin Px – FeO	1			-4.65		-		,			-							,		
WLR all	13	-22.23	-15.29	-19.56	0.23	12	-1.50	3.04	0.50	0.19	12	3.68	6.54	4.78	0.24					
WLR Mel	4	-19.84	-16.81	-18.45	0.45															
WLR OI	3	-22.23	-17.97	-20.35	0.49	1			-0.37											
WLR Px	4	-21.66	-15.29	-19.32	0.36	4	-1.22	3.04	0.86	0.25	4	4.95	6.54	5.16	0.99	8	-0.34	9.97	2.72	0.16
WLR Sp	2	-22.05	-20.88	-21.46	0.68	7	-1.5	1.25	0.19	0.32	7	3.70	5.70	4.76	0.25					
AR all	6	-22.41	-12.81	-19.00	0.35															
AR Fo	3	-22.41	-21.81	-22.18	0.49	5	-1.49	1.39	-0.02	0.25	5	0.01	0.09	0.06	0.04	4	-1.88	-0.46	-1.34	0.24
CAIs in AR [†]	3	-17.62	-12.81	-15.83	0.49															

Table 2 Summary of O, Mg and Si isotopic analyses in E101.1 lithological units and minerals.

Abbreviations, min = minimum, max = maximum, wm = weighted mean, wm $2\sigma = 2$ sigma uncertainty on the weighted mean, sub = subinclusion, sin = sinuous, Mel = melilite, Sp = spinel, Px = pyroxene, Ol = olivine, Fo = forsterite.

* Mel in close contact with sp cluster. ** Except AR Fo data given as δ^{26} Mg excess in ‰.

[†] CAI-like nodules within the AR.

[#] Number of analyses.

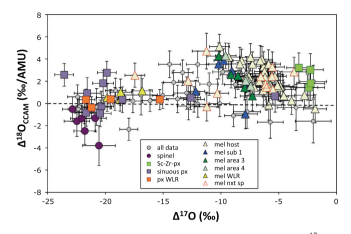


Fig. 7. Apparent oxygen isotope mass fractionation relative to the CCAM line as a function of Δ^{17} O. All data include perovskite, olivine, nepheline, FeO-rich silicates and glass. 2σ errors.

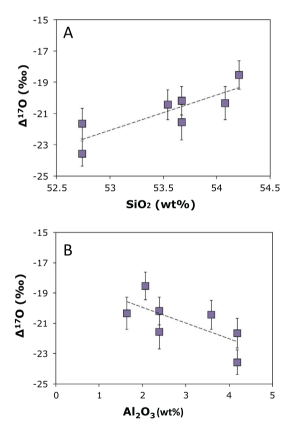


Fig. 8. Oxygen isotopic composition of sinuous pyroxene as a function of chemistry. (a) Δ^{17} O as a function of SiO₂ content. (b) Δ^{17} O as a function of Al₂O₃ content. Error bars are 2σ .

as with initial δ^{26} Mg^{*} in AOAs calculated from internal isochrons (MacPherson et al., 2012; Mishra and Chaussidon, 2014).

3.3.2. Mass fractionation

 δ^{25} Mg values vary between 6.7 \pm 1.8‰ and -4.7 \pm 0.7 (2 σ , Table S2). This ~12‰ range is unusual in normal (i.e., non-FUN) CAIs, which commonly have more

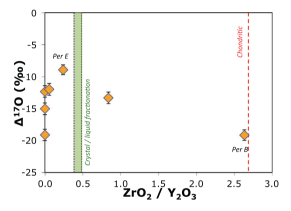


Fig. 9. Oxygen isotopic composition of perovskite as a function of chemistry. Per E and Per B refer to grains described in El Goresy et al. (2002). The chondritic ZrO_2/Y_2O_3 ratio and that expected from crystal/liquid fractionation are shown by a red dashed line and a green bar, respectively. 2σ errors. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

homogeneous compositions (e.g., Simon et al., 2005; Kita et al., 2012; Bullock et al., 2013). The Mg isotopic compositions of the E101.1 lithological units span a large range between that of light hibonite CAIs from CM chondrites (δ^{25} Mg values as low as -7%, Liu et al., 2012) and that of classical igneous Type A and B CAIs enriched in heavy isotopes ($-1\% \leq \delta^{25}$ Mg $\leq +11\%$, Davis and Richter, 2014).

The most negative values are systematically found in the sinuous pyroxene fragments with δ^{25} Mg values between -4.7 and -2.4%. In most of the inclusion, melilite has positive values between 1.9% and 6.5% with a mean value around $\sim +5\%$. Intermediate δ^{25} Mg values are found in melilite in areas 3 and 4 and in the subinclusion 1, all in close vicinity of the sinuous pyroxene ($-1.5\% \le \delta^{25}$ Mg $\le +2.1\%$). Spinels in clusters 1–4 have positive values comparable to that of the associated melilite and close to the typical host melilite value (Fig. 12, Table 2). In cluster 5, spinels have lower δ^{25} Mg values ($\sim 2.6\%$). One associated melilite has a comparable low value (3.7%) whereas

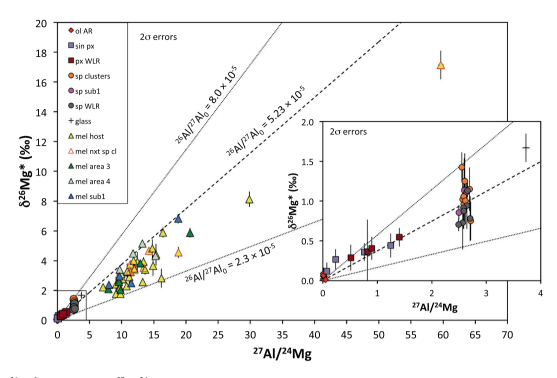


Fig. 10. δ^{26} Mg^{*} as a function of 27 Al/ 24 Mg ratio. Inset shows close-up view of low Al/Mg phases. The dashed line indicates the theoretical isochron for a canonical initial 26 Al/ 27 Al ratio and the two dotted lines show the theoretical isochrons passing through spinel with the highest initial 26 Al/ 27 Al ratio and through melilite with the lowest initial 26 Al/ 27 Al ratio. 2 σ errors.

another melilite is more typical of the host melilite (6.1‰). Both spinels and melilite in subinclusion 1 have near 0‰ $\delta^{25}Mg$ values. In the WLR, the $\delta^{25}Mg$ values of pyroxene range between -1.2% and positive values approaching those of the interior melilite at +3%. WLR spinel and forsterite from the AR all have $\delta^{25}Mg$ values around $0\pm1.5\%$.

3.4. Silicon isotopes

The δ^{30} Si values in E101.1 span a very large range, from -9.0 ± 1.3 to $+19.9 \pm 1.3$, which corresponds to FSi values between $-4.5 \pm 0.6\%$ /amu and $\pm 10.0 \pm 0.6\%$ /amu where errors correspond to 2σ uncertainties (Table S3). This $\sim 15\%$ /amu range is comparable to that of δ^{25} Mg or slightly larger. Again it is highly unusual in non-FUN CAIs, which commonly have a more homogeneous composition (Shahar and Young, 2007; Bullock et al., 2013). The lightest Si isotopic composition (most negative FSi values) corresponds to the strongest negative mass fractionation ever reported for a CAI. By contrast, the heaviest Si isotopic compositions of E101.1 reach those of the heavy (although not the heaviest) FUN CAIs (having δ^{29} Si values commonly larger than +5‰, Davis and Richter, 2014). This corresponds to a positive mass fractionation even stronger than that for Mg.

As is the case for Mg isotopes, the most negative FSi values are also found in the sinuous pyroxene fragments (-4.5 $\pm 0.6\%$ /amu to -2.4 $\pm 0.6\%$ /amu). FSi values in the host melilite are by contrast systematically positive and range

between +3.9‰/amu and +8.5‰/amu with an average and representative value at +5.9‰/amu, approaching the transition between normal and FUN CAIs (Davis and Richter, 2014). Melilite in areas in close vicinity to the sinuous pyroxene fragments (sub1, area 3 and 4) have FSi values intermediate between those of the host melilite and those of the sinuous pyroxene, although they can reach the host melilite values $(+1.2 \pm 0.4\%)/\text{amu} \le F\text{Si} \le +8.8$ \pm 0.6%/amu). AR olivines have slightly negative FSi values within 2σ errors of the average value (FSi between $-1.9 \pm$ 0.4‰/amu and -0.5 ± 0.6 ‰/amu, average $FSi = -1.2 \pm$ 0.6%/amu) and pyroxenes in the WLR have FSi values ranging between those of the AR and those of the interior melilite $(-0.3 \pm 0.3\%)/\text{amu} < FSi < +6.2 \pm 0.6\%/\text{amu}$ with one value at $\pm 10.0 \pm 0.6\%$ /amu. The latter analysis corresponds to Ti-rich pyroxene finely intergrown with spinel, interior to the Al-diopside layer of the WLR. It must be treated with caution as the analysis overlapped significantly with spinel.

3.5. Profiles

Possible isotopic variations between the interior of the CAI and its outer part were investigated with two profiles, starting from the rim toward the interior, and with the areas 3 and 4 (see Fig. 1A). Due to large ion beam spot sizes compared to the usual WLR layers, rim analyses were acquired where the thickness allowed it. No transect was done within the WLR and the profiles started inside the spinel layer without including the WLR. The chemical composition of

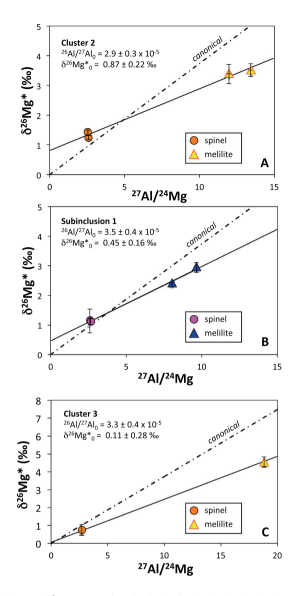


Fig. 11. Al/Mg systematics of paired spinel and melilite. (A) cluster 2. (B) subinclusion 1. (C) cluster 3. Error bars are 2σ . 2σ uncertainties on slope and intercept.

melilite shows large variations between Åk₂₀ and Åk₄₀ (profile 1, Fig. 13a) in the CAI interior, (also visible in the Mg map; Fig. 2). In the outermost 200 µm the composition of melilite is rather homogeneous at \sim Åk₂₅, except in the vicinity of the rim, where the Åk content suddenly drops towards nearly pure gehlenitic composition. This drop occurs \sim 30 µm inward from the WLR in profile 1 and \sim 70 µm inward from the rim in profile 2 (Fig. 13a). This enrichment in gehlenite toward the rim is commonly observed in Type A CAIs (e.g., Fahey et al., 1987; Goswami et al., 1994; Aléon et al., 2007; Simon et al., 2011, 2016; Katayama et al., 2012, Kawasaki et al., 2012, 2017).

The change in Mg content of melilite along profiles 1 and 2 is correlated with an isotopic evolution. Along profile 1. the δ^{25} Mg value is around +4.5% in the interior of the CAI, where melilite is the most magnesian. In the $Åk_{25}$ zone, it is rather homogeneous around +5.5‰ until the point where the Åk content drops (Fig. 13b). Unfortunately, it was not possible to acquire a Mg isotope analysis in the 30 µm region immediately inside the WLR without overlapping with spinel. However, because the point where the Åk content drops in profile 2 is further away from the rim at \sim 70 µm toward the interior, several analyses were obtained in this particular region. In the interior part of profile 2, the δ^{25} Mg value barely decreases from +6.7‰ down to + 5.1% until the 70 µm boundary (Fig. 13b). In the outer 70 μ m, it decreases from +5% down to +2% (Fig. 13b). Such an evolution from heavy, positive values in the interior toward values closer to 0 has already been observed in some Type A CAIs (Goswami et al., 1994; Simon et al., 2005), although the reverse has also been observed (e.g., CAI Leoville 144A, Simon et al., 2005). Areas 3 and 4 also show an isotopic evolution with distance but are generally much closer to 0, with δ^{25} Mg values evolving from about +2% down to $\sim0\%$ toward the rim (Fig. 13b).

This coupled chemical - stable Mg isotopic evolution of melilite is also roughly associated with a radiogenic Mg isotopic evolution. $({}^{26}\text{Al}/{}^{27}\text{Al})'_0$ ratios are scattered between $\sim 4 \times 10^{-5}$ and $\sim 2 \times 10^{-5}$ in the interior of the CAI but

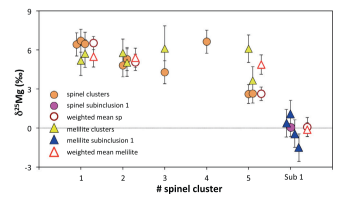


Fig. 12. δ^{25} Mg of spinel clusters and associated melilite. In subinclusion 1, only melilite data in the close vicinity of analyzed spinel are shown. 2σ errors.

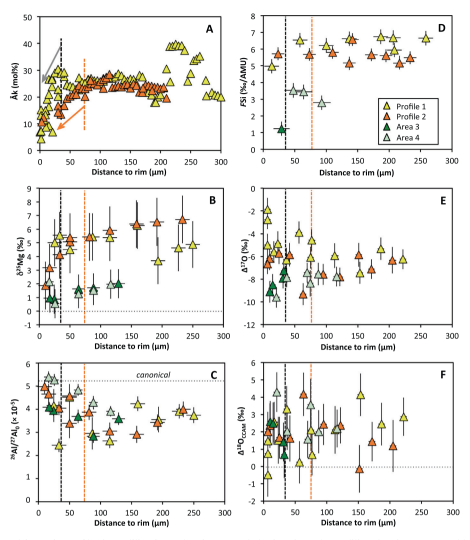


Fig. 13. Chemical and isotopic profiles in melilite from the rim toward the interior. (A) Melilite chemistry. (B) Stable Mg isotopes. (C) Inferred initial 26 Al/ 27 Al. (D) Stable Si isotopes. (E) 16 O excess. (F) O isotopic mass fractionation (deviation from the CCAM line). Black dashed line - limit of transition zone between interior and WLR along profile 1 (where melilite becomes more gehlenitic as illustrated by grey arrow), orange dashed line - limit of corresponding zone along profile 2 (melilite behavior illustrated by orange arrow). Errors on the x-axes correspond to the beam size. 2σ errors on the y-axes. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

increase toward the rim, notably outward of the 70 μ m limit in profile 2, to reach roughly canonical values near the rim (Fig. 13c) as also found by Simon et al. (2005) for CAIs with supracanonical ratios in their interior.

By contrast, the Si isotopic composition does not show any appreciable evolution with distance from the interior to the rim (Fig. 13d). Profiles 1 and 2 are flat with homogeneous FSi values typical of the interior CAI near $\sim+6\%$ / amu. Areas 3 and 4 also have seemingly homogeneous compositions but at different FSi values near + 3‰ (area 4) and + 1‰ (area 3), although only 1 value without significant overlap is available for the latter.

Similarly to Si, the O isotope profiles do not show clear trends along the profile. Δ^{17} O values along profile 1 cluster around -6% with a significant spread as discussed above. Still, the outermost two analyses show Δ^{17} O values near -2% near the rim (Fig. 13e). Profile 2 shows comparable values (Fig. 13e) but a slight decrease in Δ^{17} O is observed

toward the exterior before increasing again in the outer 70 µm. This increase and the least negative values of profile 1 may indicate a systematic increase in Δ^{17} O near the rim. Such an increase in Δ^{17} O is uncommon in Type A CAIs (e.g., Aléon et al., 2007; Kawasaki et al., 2012; Simon et al., 2016) but has been observed previously (e.g., CAI Allende A37, Simon et al., 2011, 2016). Areas 3 and 4 show significantly lower Δ^{17} O values (near -8%c, see above), again with some scatter. Contrary to profiles 1 and 2, Δ^{17} O values tend to decrease toward the exterior in area 3 and 4. The mass fractionation relative to the CCAM line is highly variable and does not show any systematic trends (Fig. 13f).

4. DISCUSSION

The coupled isotopic and petrologic observations summarized above point to a complex origin for E101.1. Here,

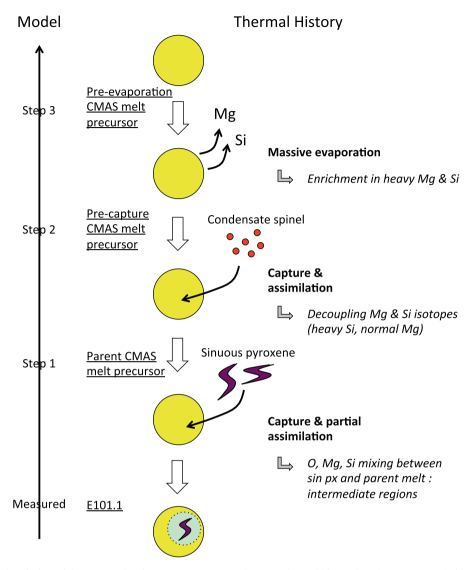


Fig. 14. Schematic description of the events taken into account to unravel E101.1 thermal history based on O, Mg and Si isotopes. The Model arrow illustrate the backwards calculations to find the precursor composition. Steps 1-3 are discussed in text. Endmember compositions are given in Table S4.

we present model calculations to determine the nature of the E101.1 precursor materials based on observed isotopic compositions. The models reported in sections 4.1 and 4.2 are depicted schematically in Fig. 14. Theoretical endmember components are listed in Table 1; their compositions are given in supplementary information (Table S4).

4.1. Origin of the sinuous pyroxene

4.1.1. Thermal history of the sinuous pyroxene

The very light Si and Mg isotopic compositions of the sinuous pyroxene fragments with δ^{25} Mg and FSi values both of about -3.5%, are unusual in igneous CAIs, which are commonly enriched in heavy isotopes due to Rayleigh distillation during evaporation (e.g., Clayton et al., 1988; Grossman et al., 2008b). These compositions are drastically different from those of the surrounding host CAI, which is enriched in heavy Si and Mg isotopes. This is indicative of a

strong degree of disequilibrium and confirms the hypothesis inferred from textural and petrographic analysis (El Goresy et al., 2002) that the sinuous pyroxene lithologies are indeed xenolithic fragments that were captured by the main host and escaped complete assimilation.

Isotopically light values of Mg have already been reported for many platy hibonite crystals (PLACS) from the Murchison and Paris CM chondrites, with δ^{25} Mg values occasionally reaching -7% and possibly as low as -10%(Fahey et al., 1987; Ireland, 1988, 1990; Sahijpal et al., 2000; Liu et al., 2012, Kööp et al. 2016a, 2016b). The proposed interpretation for such low δ^{25} Mg values is kinetic isotopic effects during condensation (Liu et al., 2012), at a temperature lower than the equilibrium condensation temperature. The competition between kinetic and equilibrium effects has been investigated theoretically for the evolution of Mg isotopic composition during condensation of forsterite (Richter, 2004) and experimentally for the evaporation of a melt of CAI composition (Richter et al., 2007). The amount of Mg isotope mass fractionation due to kinetic effects can reach -10 to -14%/amu, depending on temperature. A contribution of Mg and Si condensed under thermal disequilibrium thus appears to be a possible explanation for the negative δ^{25} Mg and FSi of the sinuous fragments.

A key parameter in the extent of the kinetic effects is the ratio between the condensation timescale and the timescale for the temperature change of the ambient medium (Richter, 2004). If the condensation is much faster than the change in ambient temperature, then condensation mostly proceeds at equilibrium and the isotopic fractionation is negligible in the considered range of temperatures. By contrast, if the change in ambient temperature is faster than condensation, then condensation proceeds at a temperature lower than expected (undercooling) and large kinetic isotopic fractionation effects are to be expected (Richter, 2004; Simon and DePaolo, 2010; Simon et al., 2017). Kinetic isotopic fractionation during condensation thus appears as a possible explanation for the negative δ^{25} Mg and FSi values of the sinuous pyroxene if condensation proceeded during a rapid temperature drop (Richter, 2004) allowing a significant degree of undercooling (Simon and DePaolo, 2010; Simon et al., 2017). Either the sinuous pyroxene contains a small fraction of a strongly fractionated component or a large fraction of a component having a few per mil fractionation (e.g. a few degrees of undercooling as calculated for Ca and Ti isotopes, Simon et al., 2017). Such a temperature drop could be due to fast removal by rapid transport of condensing precursors from a region of high temperature in the protoplanetary disk, where Mg and Si start to condense in solids, to a region where the ambient temperature is suitable for pyroxene condensation. Alternatively rapid temperature fluctuations are required.

Interestingly, the sinuous pyroxene exhibits the UR REE pattern typical of E101.1, wherein the relative abundances of REEs are controlled by their respective volatilities, the most refractory REEs being the most abundant. The UR REE pattern is established by condensation as solid solutions in the most refractory minerals that can accommodate REEs, namely ZrO₂, Y-oxides, hibonite and perovskite. As hibonite is the first major condensate, the UR pattern is established in the first few degrees below the onset of hibonite condensation (Simon et al., 1996; Davis and Richter, 2014). Such REE patterns can only be preserved in the highest temperature condensate (hibonite ?) if it is rapidly removed from the gas. One may expect a significant amount of thermal disequilibrium during this removal of the UR-carrier from the ambient gas, which possibly accounts for Ca and Ti isotopic mass fractionation (Davis et al., 2018; Simon and DePaolo, 2010; Simon et al., 2017). Whether it also accounts for the negative Mg and Si isotopic fractionation is unclear. A separate condensation event may be required due to the large difference (~ 200 K) in condensation temperatures between Ca and Ti on one hand, and Mg and Si on the other hand. Examination of UR-CAIs for which both REE patterns and Mg isotopes are available indeed shows that CAIs with UR REE patterns often have negative δ^{25} Mg values between 0 and -10%, albeit with relatively large uncertainties (Hinton et al., 1988; Fahey et al., 1994; Simon et al., 1996, 2002; Liu et al. 2009). The only known UR CAI with $\delta^{25}Mg > 0\%$ (CAI 3483–3–10 in Mighei, MacPherson and Davis, 1994) is an altered fragment with a WLR. The combination of heavy Mg isotopic composition and presence of a WLR suggests that it once was an igneous CAI having undergone evaporation.

One possible scenario to explain the REE and isotopic data is that the sinuous pyroxene formed from hiboniterich precursors that were rapidly removed from the high temperature regions where the UR REE pattern was established and then transported to a region where the temperature, about 200 K lower, was favorable for diopside condensation. The rapid transport would have resulted in preservation of the UR REE pattern at the onset of removal and in subsequent kinetic isotopic fractionation during Si and Mg condensation.

4.1.2. Partial assimilation of the sinuous pyroxene

The systematic comparison of the O, Si and Mg isotopic compositions of melilite throughout the inclusion shows that the melilites from areas 3 and 4 and from the subinclusion 1 have isotopic compositions intermediate between those of the Al-diopside sinuous pyroxene fragments $(\Delta^{17}O \sim -20\%, \delta^{25}Mg \text{ and } FSi \sim -3.5\%)$ and the remaining melilite of the host inclusion, which cluster around Δ^{17} O ~-6‰, δ^{25} Mg ~+5.5‰ FSi ~+6.5‰. These intermediate isotopic compositions can be reproduced by a single mixture between the Al-diopside sinuous pyroxene and the E101.1 main host (Fig. 15) with the exception of two spots in subinclusion 1 (Fig. 15c). The shape of the mixing hyperbolae depends on the relative concentrations of the elements, but by considering the O, Mg and Si isotope compositions we can calculate the mixing proportions in the main E101.1 host by assuming a composition for the second end-member (Fig. 14). The sinuous fragments are dominated by Al-diopside with Al₂O₃ typically in the 2-5 wt% range. For the calculation, we thus assumed that this endmember is pure diopside (CaMgSi₂O₆) having the isotopic composition of the sinuous pyroxene. If we further assume that the second end-member consists only of the five elements Ca, Al, Mg, Si and O, we find that the three mixing hyperbolae cannot be reproduced simultaneously by any known mineral having the isotopic composition of melilite. For example, the Δ^{17} O- δ^{25} Mg mixing line could be matched by mixing diopside with melilite in the Åk₁₀-Åk₃₀ range with a best match for $\sim Ak_{20}$, but even Ak_{10} would still be too Si-rich to provide a match for the Δ^{17} O-FSi mixing line. Rather, this suggests that the mixing occurred between Al-diopside and a CMAS melt having isotopic compositions equal to those observed in melilite. With the final assumption that such a melt had a solar Ca/Al ratio, as expected for most CAIs (e.g., Simon and Grossman, 2004; Grossman et al., 2008b), the data can be fit by mixing hyperbolae corresponding to an extremely refractory CMAS melt having approximately 49 wt% Al₂O₃, 39 wt% CaO, 10 wt% SiO₂ and 2 wt% MgO. This implies that the sinuous Al-diopside fragments are relicts from a xenolith mostly made of ¹⁶O-rich diopside with light Mg and Si

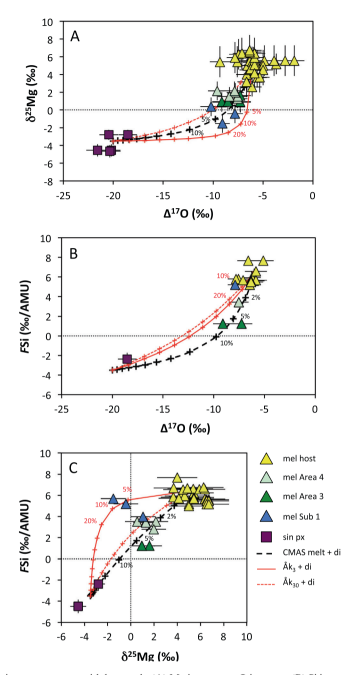


Fig. 15. Mixing calculation of sinuous pyroxene with host melt. (A) Mg isotopes vs O isotopes. (B) Si isotopes vs O isotopes. (C) Si isotopes vs Mg isotopes. Mixing lines between sinuous pyroxene (assumed to be diopside) and a second component having the isotopic composition of the typical host melilite are shown. Black dashed line: the second component is a refractory CMAS melt. Red lines: the second component is melilite of indicated composition. Mixing proportions are given. 2σ errors. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

isotopic compositions that was captured by an extensively molten, highly refractory, ¹⁶O-poor melt. They were partially assimilated in this melt, which subsequently crystallized as melilite with intermediate isotopic compositions in area 3 and 4 and in subinclusion 1, while the remaining melilite preserved the initial melt composition upon crystallization. This melt is hereafter referred to as the step 1 parental melt (Table 1). The two points in subinclusion 1 not matched by this mixing are instead well reproduced by using an endmember of almost pure gehlenitic composition. This suggests that the CMAS melt was partly crystallized as gehlenite within subinclusion 1, when the sinuous pyroxene was captured. The existence of such a partially crystallized region may explain why some sinuous pyroxene fragments distribute around subinclusion 1. The observed compositions are reproduced if 2–5 mol% of sinuous pyroxene was assimilated in areas 3 and 4, while this proportion reach 10 mol% in subinclusion 1.

Most likely all the sinuous pyroxene fragments derived from a single proto-CAI as indicated by (1) similar, unusual mineralogy and isotopic compositions and (2) the mixing lines between the sinuous fragments, areas 3 and 4, subinclusion 1 and the main host, which do not require more than one trapped xenolith. Detailed examination of the REE patterns (El Goresy et al., 2002) suggests that the sinuous pyroxene fragments are genetically related to the rest of the E101.1 CAI. Indeed, all UR CAIs have distinct REE patterns that record complexities in condensation events. These differences are seen most clearly in the abundances of light REE (LREE) relative to those of the heavy REE (HREE) and in the relative proportions of Lu, Er and Ho, the most refractory REEs. In E101.1, all units except a few Sc-Zr-rich pyroxene spots have abundant LREE relative to the HREE. In addition, Lu, Er and Ho are present in comparable abundances in E101.1. Given that these elements have different volatilities, direct condensation should result in Lu > Er > Ho (Davis and Grossman, 1979; Simon et al., 1996; Davis and Richter, 2014). Their comparable or reversed abundances are usually attributed to fractional condensation and removal of an UR mineral with uniform REE activity coefficients, such as ZrO2, that would preferentially remove Lu and then Er from the gas before establishment of the UR pattern in the main hibonite or perovskite condensate (Simon et al., 1996). The sinuous pyroxene fragments share both the LREE and Lu/Er/Ho features with the rest of E101.1 indicating that they may have been initially related with the host inclusion, perhaps by common UR precursors in spite of a totally different thermal history.

It has been suggested that these diopside-rich xenoliths could represent a former generation of WLR (El Goresv et al., 2002). However, our data do not allow a firm identification of the sinuous pyroxene precursor. Several features indeed compare well with WLRs such as its layered structure, the Al-diopside chemistry of pyroxene commonly found as the outer layer of WLRs and the common association of diopside with anorthite. It is not clear if the sinuous pyroxene is a former WLR fragment but several hypotheses can probably be discarded. The sinuous pyroxene is unlikely to be a fragment from the E101.1 WLR pushed inside because its Mg and especially Si isotopic composition are more negative than those found in the WLR of E101.1 and because the latter contains a Ti-rich fassaitic pyroxene layer inside diopside instead of anorthite (El Goresy et al., 2002). The sinuous pyroxene is also unlikely to be a remant of WLR associated with subinclusion 1. If our analysis is correct, subinclusion 1 is just a region of the host resulting from assimilation of a large fraction of sinuous pyroxene. Its rounded shape is probably fortuitous. Furthermore, in several places the sinuous pyroxene fragments around subinclusion 1 contain diopside in contact with subinclusion 1 and anorthite outside, which is the reverse of the commonly observed WLR sequence. It further lacks the spinel layer ubiquitous in WLR.

A similar approach can be used to investigate if the quenched glass areas with relict pyroxene and skeletal gehlenite (Fig. S1) derive from a secondary impact melting event, as initially proposed (El Goresy et al., 2002), or represent a residual melt from the trapping and partial

assimilation of the sinuous fragments. Here caution must be taken as (1) only one Mg isotope analysis (Mg36) and no Si isotope analysis has been performed because numerous fine-grained mineral inclusions in the glass prevented more analyses, and (2) matrix effects in Mg isotopic analyses in glass relative to stoichiometric pyroxene or melilite may result in incorrect Mg isotopic ratios. If the analysis is treated as a pyroxene glass, the mixing calculation shows that the O and Mg isotopic compositions of the quench melt regions are matched by the mixing hyperbola calculated for the refractory CMAS parental-melt described above (Fig. S6), in which case the quench glass may be a residual melt. In contrast, if a matrix effect intermediate between that of pyroxene and melilite is applied to account for the glass composition intermediate between diopside and melilite, then the mixing is more consistent with mixing the sinuous pyroxene with a melilite component $\sim Ak_{30}$ in composition (Fig. S6). In this case an impact origin is possible. Dendrites in this melt require rapid cooling, which may imply a two stage history. Note that the presence of Fe in the quenched glass (FeO up to 0.9 wt%) indicates a contribution of the FeO-rich regions associated with the sinuous pyroxene, which may have locally changed the composition of the mixture toward a less refractory and more ¹⁶O-poor composition. The presence of FeO in the glass indicates the FeO-rich regions must have predated formation of the quenched glass (see Section 4.6).

4.2. Thermal history of the main host

The main host precursor as inferred from the mixing calculation between Al-diopside and present-day host melilite is a highly refractory CMAS melt enriched in the heavy Si and Mg isotopes. The Si and Mg isotopic fractionations imply a significant amount of evaporation, but it is not clear whether this evaporation was sufficient to explain the ultrarefractory composition of the CMAS parental melt that was highly depleted in Si and Mg. The Si isotopic composition (FSi $\sim +6.5\%$) is comparable with that of several F- and FUN inclusions (e.g., TE, CG-14, EK-1-4-1, Clayton et al., 1988; Mendybaev et al., 2013) as well as that of evaporation residues produced in the conditions expected for the formation of FUN CAIs (Mendybaev et al., 2013, 2017). However, the Mg isotopic composition $(\delta^{25}Mg \sim +5\%)$ is closer to that of regular CAIs, whereas Mendybaev et al. (2013, 2017) show that δ^{25} Mg values between +10 and +15% are expected from evaporation experiments and up to +20% are found in natural FUN CAIs (e.g., CG-14, EK-1-4-1) with FSi comparable to that of E101.1, the lowest δ^{25} Mg values being from evaporation of a forsterite-rich precursor (Mendybaev et al., 2017). The larger degree of mass fractionation in Mg than in Si in these experiments is due to the evaporation of Mg as Mg and Si as SiO, which fractionates less due to its higher mass. Either the E101.1 compositions are indicative of unusual evaporation conditions or the Si and Mg isotopic fractionations were decoupled to some extent. In light of the capture and partial assimilation of the sinuous pyroxene, another capture event can possibly explain the low Mg isotopic composition if the captured material is Mg-rich and Si-poor, such as spinel, and has the usual δ^{25} Mg value expected from equilibrium condensation near 0‰ if formed from a reservoir with normal planetary Mg isotopic composition. The CMAS composition of the parent melt predating this second capture event is hereafter referred to as the step 2 precapture melt.

To understand the origin of the parent material of the E101.1 main host, we calculated what would be its preevaporation chemical composition and the extent of Si and Mg loss, based on the measured Si and Mg isotopic compositions and the CMAS composition of the step 1 parental melt calculated above. To account for possible spinel capture, we considered several cases: (1) no additional material, (2) assimilation of condensate spinel with $\delta^{25}Mg = 0\%$ so that the initial melt composition had a δ^{25} Mg of +10‰ and (3) assimilation of condensate spinel so that the initial melt composition had a $\delta^{25}Mg =$ +15%. In cases of assimilating spinel mass balance considerations indicate that the $+5\% \delta^{25}$ Mg value is achieved if half of the Mg comes from spinel (case 2) or two-thirds of Mg comes from spinel (case 3). These contributions correspond to a mass fraction of spinel vs CMAS melt of 3.6 wt% and 4.8 wt%, respectively, which is not unreasonably large. The step 2 precapture composition of the parental melt is thus calculated by removing the Al and Mg contribution of these amounts of spinel from the step 1 parental composition. Pre-evaporation calculations were subsequently done in the three cases, (1) assuming that Si and Mg evaporate as $SiO_{(g)}$ and $Mg_{(g)}$, (2) using the Rayleigh equation for kinetic evaporation and (3) using either the fractionation factors determined in Knight et al. (2009) for a sub-liquidus temperature of Type B inclusions of 1400 °C, or the fractionation factors determined by Mendybaev et al. (2013) for the production of FUN inclusions at 1900 °C, thus yielding six possible pre-evaporation compositions referred to as step 3 compositions.

The results presented in Fig. 16, show first that the contribution of captured spinel does not change significantly the composition of the parental melt (step 2) relative to the products of equilibrium condensation as calculated in Grossman et al. (2008b). In each of the three cases, it is close to the first condensate expected at high temperature, near the field of melilite + krotite (CaAl₂O₄) condensation. However, the heavy Si and Mg isotopes indicate that this cannot be the initial composition of the parental E101.1 host melt. Second, using any set of kinetic fractionation factors yields comparable results. All six step 3 pre-evaporative compositions calculated with any amount of spinel removed at step 2 are found to match nicely with the theoretical composition of a condensate (Fig. 16) in the hibonite + perovskite + melilite field at 10^{-3} to 10^{-4} bar (Mendybaev et al., 2017).

The precursor material of the E101.1 parental melt is thus consistent with an extremely refractory condensate, much more refractory than typical Type A CAIs, that subsequently underwent evaporation. In the model without spinel assimilation, the fractions of Si and Mg lost are 47% and 42%, respectively, using the fractionation factors of Knight et al. (2009) and 47% and 29%, respectively using those of Mendybaev et al. (2013). Assuming that the Mg-Si isotopic decoupling is due to condensate spinel assimilation, the fraction of Mg lost reaches 66%/50% (depending on the fractionation factor) for an initial δ^{25} Mg value of +10‰ and 80%/64% for an initial δ^{25} Mg value of +15‰.

The highly refractory composition of the precursor condensate, in the stability field of hibonite, perovskite and melilite, is well in line with the presence of large perovskite grains. It also agrees with the UR REE pattern, which points to a condensation temperature range corresponding to that of hibonite and perovskite, the highest temperature major condensates that carry REE (e.g., Lodders, 2003). The excesses and depletions in ⁴⁸Ca ($-7.5\% \le \delta^{48}$ Ca \le

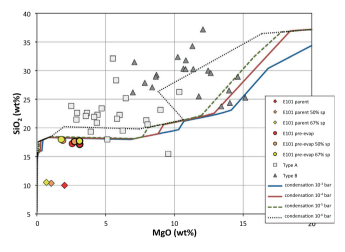


Fig. 16. Comparison between the E101.1 parental CMAS melt composition (diamonds) and its pre-evaporation composition (circles) with the expectation of the equilibrium condensation sequence (trajectories from Mendybaev et al., 2017) and the composition of regular type A CAIs (open squares) and B CAIs (grey triangles). Red symbols - no Mg considered to come from subsequent spinel assimilation, orange - 50% Mg considered to come from subsequent spinel assimilation. Bold circles: calculated with fractionation factors from Knight et al. (2009). Thin circles: calculated with fractionation factors from Mendybaev et al. (2013). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

+9.8%) in three perovskite grains analyzed previously (El Goresy et al., 2002) suggests a possible connection with FUN inclusions, agreeing with the strong mass fractionation by evaporation indicated by the FSi values. Although unusual, such ⁴⁸Ca anomalies without associated ⁵⁰Ti anomalies have also been found in hibonite-rich FUN inclusions from Murchison (Kööp et al., 2018). Together with the calculated hibonite-rich precursor composition and the UR pattern, often attributed to hibonite condensation (e.g. Davis et al., 2018), this suggests a link between the precursor of E101.1 and hibonite-rich FUN inclusions. Whether the fractionated component (F) is initially the same as the carrier of the nuclear anomalies in ⁴⁸Ca in E101.1 is unclear because the subsequent capture of xenolithic material renders the characterization of the initial material subject to uncertainties. The precursor material of E101.1 can be viewed as an ultrarefractory FUN inclusion resulting from extensive melting and evaporation of an UR condensate precursor with isotopic anomalies of nucleosynthetic origin. Alternatively, the UR component with nuclear anomalies (the UN component) recorded in the pristine condensate perovskite may have been aggregated to a highly mass fractionated component (the F component) before they were partially melted together. In this case the F and UN components could have been initially decoupled as suggested by the existence of F CAIs without nuclear isotopic anomalies (Clayton et al., 1984; Caillet Komorowski et al., 2007; Krot et al., 2014). The lack of large mass fractionation in O isotopes in perovskite may support this conclusion. However, many FUN inclusions contain unfractionated components issued from the first crystallization stages before the melt was completely evaporated (Thrane et al., 2008; Krot et al., 2014), which may well be the case for perovskite.

Recent studies indicate that FUN CAIs are related to normal CAIs (Park et al., 2014; Krot et al., 2014). On the one hand, Ca, Ti and ²⁶Al isotopic anomalies define a continuum between FUN inclusions and normal CAIs (Park et al., 2014, 2017). On the other hand, the coupled petrographic and O isotope study of twelve FUN inclusions shows that they are petrographically similar to normal CAIs and exhibit O isotope mass fractionation effects of variable amplitude not correlated with the isotopic anomalies (Krot et al., 2014). This was attributed to variable conditions of partial melting and evaporation and variable sampling of reservoirs with diverse presolar components (Krot et al., 2014; Park et al., 2017). If this is correct, and in light of the multiple aggregation history of E101.1 revealed by the present study, which probably includes assimilation of a FUN component, it is reasonable to consider that most igneous CAIs may have been formed by the aggregation and partial melting of various precursors, among which were a proportion of FUN material with inherited presolar isotopic anomalies.

4.3. Insights on the nature of the UR precursor of E101.1 from perovskite

Differences in the chemistries of perovskite grains in E101.1, notably in their ZrO_2/Y_2O_3 ratios, have been

interpreted as indicating the presence of four distinct populations of perovskite (El Goresy et al., 2002). These populations include perovskite grains with (1) chondritic ZrO_2/Y_2O_3 ratios ~2.7, suggesting an origin by condensation and (2) grains with ZrO_2/Y_2O_3 ratios ~0.4, typical of crystal-liquid fractionation indicative of crystallization from a melt. The two other perovskite grains are either (3) devoid of ZrO_2 or (4) have intermediate ZrO_2/Y_2O_3 ratios between 0 and \sim 2.7. Different perovskite populations can also be recognized based on their textural context, although it is not clear if chemical and textural populations can be linked: large anhedral perovksite grains are commonly associated with metal and/or V-Fe-oxide indicative of interactions with metal, and are rimmed by the Sc-Zrrich pyroxene (Fig. 4). Such textures are suggestive of partial melting and reaction of pre-existing perovskite with a silicate melt and agree with considerations from minor and trace elements (El Goresy et al., 2002). Other grains, usually much smaller, are euhedral and enclosed within pyroxene, sometimes within pyroxene \pm spinel rims around spinel (e.g., in cluster 5, Fig. 3d), suggesting late crystallization either from a Ti-rich partial melt or at the solid state. The abundance of perovskite in E101.1 suggests a perovskite-rich UR precursor of bulk chemistry in the hibonite + perovskite + melilite condensation field The chemical and textural diversity of perovskite further indicates contribution of perovskite with (1) diverse condensation histories and (2) variable post-aggregation histories including reaction with a partial melt, new crystallization from a partial melt and solid state equilibration with the surrounding phases.

Perovskite grains analyzed for O isotopes are large anhedral grains and have a range of ZrO₂/Y₂O₃ ratios. This includes (i) chondritic perovskite (per B), (ii) Zr-free perovskite (4 grains) and (iii) perovskite with intermediate ratios (2 grains). Large variations in Δ^{17} O are observed. On one hand, some grains enriched in ¹⁶O to a level approaching that of spinel ($\Delta^{17}O = -19.2\%$) are found in different populations (Fig. 9). On the other hand, both ¹⁶O-rich grains and ¹⁶O-depleted grains with Δ^{17} O up to -8.9% are found in a single population (Zr-free perovskite). Because (i) the Sc-Zr-rich pyroxene that rims perovskite is among the most ¹⁶O-poor mineral in E101.1 and most likely records the composition of the melt, and because (ii) the diffusivity of O in solid perovskite is rapid (Gautason and Muehlenbachs, 1993; Sakaguchi and Haneda, 1996), a plausible interpretation is that all perovskite grains were initially ¹⁶O-rich and experienced partial re-equilibration with the enclosing Sc-Zr-pyroxene, either sub-solidus or between solid perovskite and melt. All grains analyzed here have comparable sizes in the 20-30 µm size range and all analyses include most of the grains, so that it has not been possible to evaluate the extent of O isotopic zoning in perovskite, thus rendering the latter hypothesis impossible to confirm or negate.

The anhedral shapes of the ¹⁶O-rich perovskite grains are consistent with the idea that they represent partially melted relict grains. These grains belong to the population with close-to-chondritic ZrO_2/Y_2O_3 ratios (Per B) and to the population with ZrO_2 content near or below the detection limit of the electron probe (Fig. 9). Whereas the chondritic population can be understood as resulting from direct condensation from a solar gas, the origin of the Zr-free grains remains unclear. Because Zr-oxides have already been observed in UR CAIs (Ivanova et al., 2012) and because Zr is an ultrarefractory element that condenses at equilibrium as ZrO₂ before corundum and hibonite condense (i.e., at higher temperature, Lodders, 2003), a possibility is that the Zr-free perovskite is an UR condensate that formed after a fractionation due to previous condensation of ZrO₂. As mentioned above such a fractional condensation is in line with the comparable abundances of Lu, Er and Ho in E101.1 instead of Lu > Er > Ho as expected from relative volatilities (Simon et al., 1996). Both the chondritic and the Zr-free perovskite probably represent end-member products of the condensation of UR minerals due to slight variations of the condensation conditions (temperature or pressure). We also note that some of the Zr depletion in perovskite probably arose from the late solidstate diffusion into the Sc-Zr-rich pyroxene (El Goresy et al., 2002). A reasonable interpretation is thus that the large perovskite grains were initially part of a ¹⁶O-rich UR inclusion (or several) condensing in the hibonite + perovskite + melilite field and containing precursor UR oxides inherited from condensation steps above 1700 K (e.g., Simon et al., 1996), such as those described in other compound CAIs (e.g., Ivanova et al., 2012). Such an UR inclusion could be the carrier of the UN isotopic anomalies suggested by ⁴⁸Ca excesses and deficits in perovskite. It may have experienced a ¹⁶O-depletion during the melting event associated with extensive Mg, Si evaporation and isotopic fractionation that transformed the E101.1 precursor into a FUN inclusion. In this case, the ¹⁶O-rich perovskite would represent the starting material of the FUN inclusion and melilite $\pm\,Sc\text{-}Zr\text{-}rich$ pyroxene would represent the final crystallization product after equilibration with the ¹⁶O-poor gas (Krot et al., 2014). The lack of strongly mass fractionated O isotope component corresponding to the heavy Si component would then be explained by the peculiar chemistry of E101.1, which lacks intermediate minerals with slow oxygen diffusivities recording the composition of the evaporating melt, such as spinel, forsterite or pyroxene. The O isotopic compositions of initial perovskite and final melilite + Sc-Zr-rich pyroxene compare well with those of initial spinel and final melilite, respectively, in some FUN inclusions having Si isotopic compositions similar to that of the E101.1 parental melt (e.g., CG14, EK1-4-1, Krot et al., 2014). This scenario would also agree with the residual O isotope mass fractionation observed in Sc-Zr-rich pyroxene and some of the ¹⁶O-poor melilite (Fig. 7). Alternatively, the ¹⁶O-rich UR perovskite-rich precursor could have coagulated with a F(UN)-like proto-CAI previously mass fractionated and both were partially melted together.

4.4. Extraneous origin of the spinel clusters

Beside evidence of perovskite partial melting, there are numerous examples of spinel resorption (e.g., rounded shapes, embayments) in E101.1. These features, as well as the presence of the pyroxene-rich symplectite around spinels, and the chemical and isotopic zonation of melilite (enriched in gehlenite and often in ¹⁶O at the contact with spinel; Fig. 6) are indicative of a coronitic reaction resulting in spinel dissolution. Interestingly, a spinel + liquid = anorthite + melilite + clinopyroxene reaction has been observed in the CMAS system (O'Hara and Biggar, 1969; Schairer and Yoder, 1969; Yang et al., 1972; Onuma and Kimura, 1978). Although it has not been observed directly, such a reaction has been inferred during closed-system crystallization of Type B CAI liquids, based on the proportions of spinel as a function of temperature (Stolper, 1982). A similar reaction in the highly silica-poor system of Type A CAIs may therefore result in a comparable reaction where anorthite is absent in favor of highly gehlenitic melilite and highly aluminous pyroxene (reaction (1)).

spinel + liquid = gehlenite + clinopyroxene (1)

We suggest that the coronitic textures observed around spinel are the signature of this reaction. It would explain all resorption features in spinel, the presence of the pyroxene \pm spinel symplectite and the coupled enrichment in ¹⁶O and Al₂O₃ in melilite at the contact with spinel (Fig. 6).

Whether reaction (1) occurred in closed or open-systems cannot be deduced from the coronitic textures alone. Additional insights come from Mg isotopes (Figs. 11 and 12). Most spinel clusters have δ^{25} Mg values near 5–6‰, close to each other and within error of the associated melilite (Fig. 12). However some heterogeneities point to incomplete homogenization. Although the exact spinel-melilite difference can possibly be subject to small uncorrected matrix effects, the range of δ^{25} Mg values in spinel from 2.6% (cluster 5) to 6.6% (cluster 4) with an intermediate 4.3% (cluster 3) is beyond 2σ analytical uncertainties (<1%). The Al/Mg systematics of spinel clusters and associated melilite are mostly comparable from cluster to cluster (Table 2, Fig. 11) but heterogeneities exist as well. Intercepts in the Al/Mg regressions from the various clusters (Fig. 11) yield a range from $0.11 \pm 0.28\%$ to $0.87 \pm$ 0.22‰, again beyond 2σ errors. Finally, spinel clusters from subinclusion 1 have Mg isotope systematics similar to the others in terms of radiogenic Mg and spinelmelilite stable isotope equilibration, albeit for a much different δ^{25} Mg value near 0% (Fig. 12), which has been shown to result from mixing between the host inclusion and the sinuous pyroxene fragments (Fig. 15).

In a first interpretation, most melilite and spinel have identical Mg isotopic composition and Al/Mg systematics. In this hypothesis, supracanonical spinel in cluster 2 (Table 2, Fig. 11) and light melilite near cluster 5 (analysis Mg58) must be considered special as well as subinclusion 1, possibly another xenolith of its own in this case. Our favorite interpretation is however that these heterogeneities result from incomplete homogenization between spinel clusters and the host melilite/melt, including in subinclusion 1, where the analyzed spinel seem to belong both to the spinel swarms and to subinclusion 1 (Figs. 1 and 2). In this hypothesis, the supracanonical spinel in cluster 2 results from diffusive exchange with surrounding melilite (see Section 4.6) and the light δ^{25} Mg value at 3.7‰ near cluster 5 (Mg58) may indicate a contribution of light Mg from spinel, in agreement with O isotopes indicating a contribution of spinel-derived ¹⁶O-rich oxygen (Fig. 6).

We therefore suggest that all spinel clusters initially had a stable Mg isotopic composition different from that of the host E101.1 CAI, and thus a different origin. They subsequently underwent Mg isotope exchange with the surrounding melilite or parental melt owing to rapid Mg self-diffusion in spinel (Sheng et al., 1992) up to the point of complete or near-complete equilibration (clusters 1, 2 and 4). At 1300–1400 °C, typical of melilite crystallization, Mg isotope exchange in a 10 µm spinel is complete in 2–10 h. This isotopic disequilibrium is in line with chemical evidence for several spinel populations with varying V, Ti and Cr content (El Goresy et al., 2002). For comparable Cr content, host spinels (i.e., spinel clusters) and some spinels in subinclusion 1 have the highest V content. Because the diffusion of V in spinel is rapid compared to that of Cr (Connolly and Burnett, 2003), this indicates a significant degree of equilibration with the host melt or melilite for these spinels.

Taken together, the coronitic textures, the distribution of gehlenitic melilite and the O and Mg isotope systematics suggest introduction of the parental material of the spinel clusters by capture of solid material into a partially molten CAI host droplet, confirming the suggestion of El Goresy et al. (2002) based on the minor element chemistry of spinel. Spinel clusters can thus be considered to represent ¹⁶O-rich, spinel-rich proto-CAIs captured by the host inclusion and partially dissolved into the host melt. It is interesting to note that cluster 3, the least equilibrated of the upper 4 clusters (Fig. 1) is also the outermost. Clusters 1–4 may have been fragments of the same initial proto-CAI captured in a single event and increasingly equilibrated with melilite as they are deep within the channel.

Such a ¹⁶O-rich spinel-rich proto-CAI may have been a fine-grained-spinel-rich CAI (FG-sp-rich CAI, e.g., Krot et al., 2004a) in which minerals other than spinel, i.e., having lower melting temperature (e.g., pyroxene \pm melilite \pm anorthite), would have been entirely dissolved in the host melt, possibly contributing to an additional local enrichment in Al and ¹⁶O. In this case, the subhedral crystal shapes observed in many grains of the spinel clusters would have to result from recrystallization or annealing of finegrained aggregates of spinel. Unaltered FG-sp-rich CAIs are homogeneously rich in ¹⁶O (e.g., Aléon et al., 2002, 2005; Fagan et al., 2004; Bodénan et al., 2014; Ushikubo et al., 2017) and are commonly devoid of heavy Mg because they escaped extensive melting and evaporation. Their δ^{25} Mg values are usually near 0% or slightly negative (e.g., Clayton et al., 1988; MacPherson et al., 2010; Ushikubo et al., 2017). As a result, capture of such spinel-rich material is consistent with the Mg-Si isotopic decoupling inferred in the parental CMAS melt of the host inclusion (Section 4.2), as well as the lower δ^{25} Mg value of cluster 5. We calculated that the contribution of the spinelrich material should be of a few wt% to account for the Mg-Si isotopic decoupling, which agrees with the abundance of spinel as visible on the E101.1 polished section. Removing the spinel clusters from the host CAI mineralogy leaves an initial precursor with melilite + perovskite mineralogy,

essentially devoid of spinel, which agrees with the calculated precursor chemistry falling in the hibonite + perovskite + melilite domain if hibonite has been completely resorbed in favor of a more gehlenitic melilite. We conclude that the capture of extraneous fine-grained spinel-rich proto-CAIs well accounts for the properties of the spinel clusters as well as the isotopic systematics of the host parental melt.

The capture of ¹⁶O-rich spinel-rich CAIs by a ¹⁶O-poor melilite dominated host inclusion is analogous to the situation in E49, another CTA compound inclusion from Efremovka (Aléon et al., 2007). In other CAIs, spinel and melilite have been shown to occasionally be in Mg isotopic equilibrium (e.g., Fahey et al., 1987; Kita et al., 2012) and in some cases show evidence of Mg isotope disequilibrium (e.g., Paque et al., 2013; Goswami et al., 1994), with spinel being either heavier or lighter than coexisting melilite by at most 1 or 2‰. With the caveat that matrix effects are important in Mg isotopic analysis by SIMS and need to be carefully taken into account to investigate the spinelmelilite isotopic difference at the sub-permil level (e.g., Kita et al., 2012; Paque et al., 2013), this suggests that spinel of extraneous origin that underwent subsequent partial to complete isotopic re-equilibration may be common in CAIs.

4.5. Formation of the margins and rims of the host CAI

4.5.1. Forsterite-rich accretionary rim

The multi-isotope study of E101.1 can also be used to shed light on the origin of the outermost envelopes of igneous CAIs, the WLR and the forsterite-rich accretionary rims (AR, e.g., Krot et al., 2001). ARs are thought to be somewhat analogous to AOAs. They are composed primarily of highly magnesian olivine aggregates, commonly anhedral and porous. Some are more compact with triple junctions suggesting secondary annealing (Krot et al., 2001, 2004b). Numerous CAI-like nodules mostly composed of Ca-clinopyroxene \pm anorthite \pm spinel are often interspersed with forsterite (Krot et al., 2004b). The commonly held view is that AR are aggregates of fine-grained nebular condensates of relatively low temperature, in the range of forsterite condensation (Krot et al., 2001), subsequently aggregated onto larger CAIs and eventually partially reprocessed on the parent-body (Krot et al., 2001). The Mg and Si isotopic compositions of forsterite from the AR in E101.1 correspond to δ^{25} Mg and FSi close to 0% or slightly negative, which is in good agreement with the expectation for equilibrium condensation (Fig. 17) and with previously published Mg isotopic analyses of AR and AOAs (Larsen et al., 2011; MacPherson et al., 2012; Mishra and Chaussidon, 2014). The ¹⁶O-rich composition of forsterite indicates that the gas from which they condensed was ¹⁶O-rich as inferred from other AR and AOAs (Krot et al., 2002).

4.5.2. Wark-Lovering rim

In contrast to the ARs, the origin of WLR is more debated. They typically consist of close to monomineralic layers mimicking the condensation sequence from spinel

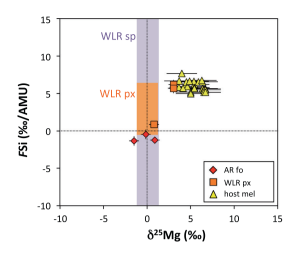


Fig. 17. Stable Mg and Si isotopic compositions of the rims on E101.1. Orange box - range of Si and Mg isotopes of WLR pyroxene (decoupled analyses), purple band - range of Mg isotopes in WLR spinel. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

 \pm hibonite \pm perovksite at the contact with the host inclusions to Al-Ti-pyroxene \pm melilite \pm anorthite, to Aldiopside (e.g., MacPherson, 2014) toward the edge of the CAI. In some cases, the outermost lavers consist of pure diopside and/or forsterite. Different origins have been proposed, which can be grouped into two categories: condensation models and flash heating models. Condensation models include (1) successive deposition onto a CAI of various layers of condensation (Wark and Lovering, 1977) with decreasing temperature from the innermost spinel \pm hibonite \pm perovskite to the outermost forsterite layers, (2) growth of isolated nucleus islands deposited onto the interior CAI (Bolser et al., 2016) and (3) diffusive equilibration of the edge of the CAI with the gas (Ruzicka, 1997; Simon et al., 2005; Simon et al., 2016). Light stable Mg isotopic composition supports these condensation models (Taylor et al., 2004; Cosarinsky et al., 2005a,b; Simon et al., 2005) but enhanced REE abundances in the rims and REE patterns identical to the host inclusion favor flash heating hypotheses (e.g. Wark and Boynton, 2001). These conflicting interpretations could potentially be reconciled if WLR formed by condensation of nebular gas and a melted outer layer is involved at some stage (Toppani et al., 2006; Keller et al., 2013; Simon et al., 2016). Mg isotopes and Ti valence have been used to argue that the WLR formed at a much higher fO₂, closer to that of chondrites, than the inner host formed at a solar fO₂ (Simon et al., 2005; Dyl et al., 2011) although this is debated (Simon et al., 2007). Finally, Mg isotopes also suggest that WLR have crystallization ages commonly younger than their host inclusions by about 200,000-300,000 years (Taylor et al., 2004; Cosarinsky et al., 2005a,b; Simon et al., 2005). Even larger time intervals of at least 590,000 and 690,000 years have recently been reported (Mane et al., 2015)

In E101.1, the outermost WLR pyroxenes exhibit the UR-REE pattern typical of the host inclusion (El Goresy

et al., 2002), including the specific LREE/HREE and Lu/ Er/Ho relative proportions, which suggests a contribution from the interior inclusion. In one case (Mg34) the δ^{25} Mg value of the WLR pyroxene of E101.1 shows an enrichment of +3%, close to that of interior melilite, but in all other cases, both pyroxene and spinel have δ^{25} Mg values near 0%, similar to those of the AR forsterite. By contrast, the FSi values span the whole range between the slightly negative values of the AR forsterite and those of the average main host melilite (Fig. 17), possibly up to $\pm 10\%$. The Mg isotopic composition is in agreement with expectations from equilibrium condensation in most cases but in the case of spot Mg34 also shows the contribution from interior Mg. The Si isotopic composition indicates a more balanced mixture between Si of condensation origin, similar to Si in the AR forsterite, and Si from the interior with enrichment in the heavy isotopes due to evaporation. Both the REE patterns and the stable Mg, Si isotopic compositions thus indicate some contribution from the interior CAI. Such an influence strongly hints that the WLR did not form by successive deposition of condensate layers at decreasing temperatures or grew from isolated nucleus islands disconnected from the interior inclusion. Nevertheless, both Mg and Si do indicate a contribution of a condensate component. Although analysis Si15 in the innermost WLR pyroxene must be treated with caution, its FSi value of +10‰, heavier than in interior melilite, may be the additional signature of a strong evaporation from a melt. Thus our data support models based on open system behavior in which the WLR formed by a combination of (i) melting of the inner host that preserved the most refractory elements and a fraction of the less refractory Mg and Si and (ii) condensation into the melt of nebular Mg and Si. Such a scenario has already been proposed from the TEM study of a WLR from a Type B1 CAI from Vigarano (Toppani et al., 2006). The latter study also documented secondary alteration phases at the sub-µm scale. Although the Nano-SIMS study of WLR from CV chondrites shows variations of ¹⁶O content (Simon et al., 2011, 2016), the study of WLR in unaltered chondrites favors a parent-body origin for these variations (Bodénan et al., 2014; Krot et al., 2017), in agreement with the TEM observations. Our data favor a systematic ¹⁶O-rich composition of primordial unaltered WLR (this work, Yoshitake et al., 2005; Aléon et al., 2007; Bodénan et al., 2014; Kawasaki et al., 2017; Krot et al., 2017). Most likely, the gas with which the CAI equilibrated during WLR formation was ¹⁶O-rich.

4.5.3. Zoning of the outer portion of the host CAI

This scenario of WLR formation explains the isotopic composition of the rim, and also the O, Mg and Si isotopic zoning observed in the outer part of CAIs (e.g., Simon et al. (2016) for O). In E101.1, chemical and isotopic profiles indicate that the distribution of Mg in the outer part of the CAI (Fig. 13) was affected by a process resulting in (1) an enrichment in Al or loss in Mg toward the exterior, (2) an enrichment in unfractionated Mg, (3) an enrichment in undisturbed near-canonical ²⁶Al. In the framework of the WLR formation by condensation from nebular gas into a partially melted CAI layer, these profiles are well

understood by a global loss of interior Mg in the periphery together with a progressive enrichment in nebular Mg. Whether the observed profiles result from solid-state diffusion, partial melting and recrystallization or a combination of both is unclear. By contrast, the Si and O isotopic compositions do not show any relationship with proximity to the WLR (Fig. 13d and e). In the case of Si isotopes, this agrees with the WLR systematics, which shows a much greater contribution of interior Si than of interior Mg in the rim (Fig. 17). Interestingly, the width of the zone affected by this interaction varies between profile 1 (30-50 µm) and profile 2 (70–100 µm) (Fig. 13a–c) suggesting small variations in the thermal regime or in the CAI response to the WLR-forming event. Parameters such as temperature, chemical gradients due to differences in total and partial pressures in the gas and to differences in CAI chemistry probably induce even larger variations of thickness of this zone between different CAIs with different thermal histories. An extreme case of intra-CAI variations may be bowl-shaped CAIs with different WLR mineralogy in their concave and convex faces suggesting different degrees of heating attributed to a lack of spinning in the nebula (Ivanova et al., 2014). Inter-CAI variations of thermal regime during WLR formation probably also explain why Mg and O gradients are not always present simultaneously. Many Type A CAIs show for instance an enrichment in gehlenite toward the exterior, in some cases associated with a Mg isotopic composition evolving toward lower δ^{25} Mg values (Fahey et al., 1987; Goswami et al., 1994; Simon et al., 2005; Cosarinsky et al., 2006), in some cases associated with an enrichment in ¹⁶O (Aléon et al., 2007; Katavama et al., 2012; Kawasaki et al., 2012, 2017). Similarly to E101.1, a decrease in δ^{25} Mg values together with little variation in Si isotopic composition has been observed in the outer mantles of Type B1 CAIs (Bullock et al., 2013). Bullock et al. (2013) conclude that these profiles would be best explained by a secondary "ad-hoc" process. We suggest that this process is in fact WLR formation. In the cases where the interior CAI is almost unaffected by WLR formation (i.e., where the transition region has a thickness near 0), the CAIs may have preserved pre-WLR zoning reflecting high temperature interaction of a solid CAI with nebular gas such as sublimation of Mg and Si from the solid state (e.g., CAI Leoville 144A, Simon et al., 2005; Shahar and Young 2007) or diffusion of ¹⁶O-poor oxygen from the gas (e.g., CAI Allende A37, Simon et al., 2011, 2016). Given the decoupling between O and Mg in E101.1, the two outermost ¹⁶O-poor analyses in profile 1 may reflect the preservation of such an initial ¹⁶O-poor composition of the melt due to equilibration with ¹⁶O-poor gas before trapping of the xenoliths and WLR formation.

4.6. Al/Mg chronology and disturbances

The lack of a well-defined Al/Mg isochron in E101.1 host melilite, the evidence of isotopic disequilibrium between spinels and host melilite, and the insufficient precision on the slopes and intercept of isochrons deduced from analyses of pyroxene preclude a simple chronological analysis of E101.1 crystallization. Still, the distribution of model $({}^{26}\text{Al}/{}^{27}\text{Al})'_0$ ratios with location in the CAI or between subcomponents can shed light on the origin of Al/Mg systematics perturbations and on the putative initial ${}^{26}\text{Al}/{}^{27}\text{Al}$ ratio prior to these perturbations (Table 2).

As mentioned in Section 4.4, the lack of complete stable Mg isotope equilibrium between spinel clusters and adjacent melilite is visible to some extent with the radiogenic isotopes. The Al/Mg systematics in paired spinel and melilite are illustrated in Fig. 11. Three cases can be distinguished.

In cluster 2 (Fig. 11a), sub-canonical mellite is associated with supercanonical spinel. All 4 analyses define a single linear relationship, which yields an initial 26 Al/ 27 Al slope of $(2.9 \pm 0.3) \times 10^{-5}$ and a distinctly positive initial ${}^{\delta^{26}}$ Mg^{*} intercept of $0.87 \pm 0.22\%$ (2σ errors). This behavior is characteristic of a closed-system redistribution of Mg isotopes between spinel and associated mellilite, pivoting around the bulk composition which would be coincident with the canonical isochron if the system initially had 26 Al/ 27 Al = 5.2×10^{-5} .

In subinclusion 1 (Fig. 11b), spinel has a canonical abundance of ²⁶Mg*, but the two melilite analyses in the vicinity of spinel define a linear array with the spinel with a subcanonical slope of $(3.5 \pm 0.4) \times 10^{-5}$ and a positive intercept of $0.45 \pm 0.16\%$. Such slope and intercepts are also indicative of closed-system exchange in the spinelmelilite system, although spinel seems to have been little affected by this perturbation. One possible explanation is an initially sub-canonical bulk composition. This would indicate a younger crystallization age for subinclusion 1, as young as 400 ± 100 thousand years after canonical. However, when considering all melilite analyses in subinclusion 1, the canonical $({}^{26}\text{Al}/{}^{27}\text{Al})_0'$ ratio of analysis Mg50 is not in a favor of such a chronological interpretation (Fig. 10). Rather it may indicate the variable contribution of a ²⁶Al-poor component (see below).

In contrast, spinel and the associated melilite of cluster 3 (Fig. 11c) define a linear array with a slope of $(3.3 \pm 0.4) \times$ 10^{-5} that passes through the origin (intercept = 0.11 \pm 0.28%). If this 2-point line is considered to be an isochron, it would yield a crystallization age of 480 ± 120 thousand years after canonical. Such an age may represent the age of cluster 3 trapping in the host CAI, however it would not be consistent with the closed-system interactions in other spinel clusters - melilite pairs implying that either the trapping of cluster 3 is a late isolated event or that the linear correlation does not correspond to the crystallization age. Another possible interpretation for this cluster closest to the edge of the CAI is an open-system reequilibration of both spinel and associated melilite with a reservoir of lower ²⁶Al content (e.g., Simon and Young, 2011). This reservoir is unlikely to be the meteorite matrix as near canonical $({}^{26}\text{Al}/{}^{27}\text{Al})_0'$ ratios are observed in melilite in close contact with the WLR in profile 1 and in areas 3 and 4 (Fig. 13c). Thus, it has to be exchange with a nebular reservoir of low ²⁶Al content.

The distribution of model $({}^{26}\text{Al}/{}^{27}\text{Al})'_0$ ratios in melilite as function of distance from the rim (Fig. 13c) is helpful in interpreting the overall Al/Mg isotope systematics of E101.1. As mentioned above, profile 2 and areas 3 and 4 display near canonical $({}^{26}\text{Al}/{}^{27}\text{Al})_0'$ ratios in the vicinity of the rim. Apart from one analysis with a low $({}^{26}\text{Al}/{}^{27}\text{Al})_0'$ ratio in profile 1, a progressive decrease in $({}^{26}\text{Al}/{}^{27}\text{Al})_0'$ ratios toward the interior is visible everywhere in the outermost 80 µm of the CAI, down to about 3 × 10⁻⁵, inward of which scattered ratios are found. Interestingly, the isotopic composition of cluster 3 is consistent with this lower limit, cluster 3 being located ~100 µm inside the CAI. As the outer part of the CAI and the WLR cannot be older than the interior CAI, this suggests a secondary perturbation with the WLR and the outermost melilite hav-

A possible interpretation based on the relationship between the parental E101.1 melt and FUN inclusions discussed above is that E101.1 crystallized from different proto-CAIs with different initial ²⁶Al contents. Indeed, many FUN inclusions are known to be devoid of or depleted in ²⁶Al (e.g., Esat et al., 1978; Clayton et al., 1984; Davis et al., 1991; Holst et al., 2013; Park et al., 2017) and mixing of a FUN precursor of low ²⁶Al content with xenoliths having canonical or near canonical ²⁶Al content will induce strong confusion of the Al/Mg isotope systematics in the final E101.1 CAI. These perturbations are likely to be more significant in the interior of the CAI than on the exterior, where re-equilibration with the regular canonical CAI reservoir, from which xenoliths are issued, will be favored. Such a ²⁶Al-poor FUN component may be the nebular reservoir required to explain the isotopic composition of cluster 3.

ing best preserved the initial near-canonical composition.

To summarize, the Al/Mg isotope systematics of E101.1 can be understood as resulting from the mixing of parental FUN material depleted in ²⁶Al with xenoliths having abundant (the sinuous pyroxene) up to canonical (the spinel clusters) ²⁶Al content. In the hot CAI interior, local closed-system interactions between spinel/pyroxene and melilite resulted in scattered (²⁶Al/²⁷Al)₀ ratios. Closer to the outside of the CAI, open-system resetting of Mg (e.g. Simon and Young, 2011) due to mixing of canonical and ²⁶Al-poor material led to intermediate Al/Mg apparent isochrons resulting from the subsequent decay of ²⁶Al (cluster 3). Finally, the outermost portion of the CAI equilibrated with the ambient gas having canonical or near-canonical ²⁶Al content. This is consistent with the initial $\delta^{26}Mg^*$ of forsterite in the AR being comparable to that determined from AOAs and AR of igneous CAIs with canonical ²⁶Al/²⁷Al ratios (Larsen et al., 2011; MacPherson et al., 2012; Mishra and Chaussidon, 2014).

A chronological constraint on the formation of E101.1 could in principle be obtained from its WLR formation, the last melting event suffered by the CAI. However, time intervals as small as $2-3 \times 10^5$ years between the formation of the interior CAI and that of the WLR (Simon et al., 2005; Taylor et al., 2005a,b; Cosarinsky et al., 2006) must be treated with caution. A problem is that the required high precision Al/Mg dating of individual minerals in WLR is extremely difficult to achieve due to the small size of the crystals. Large errors and uncertainties on the Mg isotopic closure for instance due to Mg self-diffusion in spinel make

it difficult to obtain good chronological constraints. Here we show that the range of model initial ²⁶Al/²⁷Al in WLR spinel from E101.1 varies between $(3.7 \pm 0.5) \times 10^{-5}$ and $(5.7 \pm 0.4) \times 10^{-5}$, i.e., corresponding to an apparent age interval of 460,000 years between the first and last spinel crystallized, although it is within error consistent with a 240,000 years interval between the last and the canonical spinels (Fig. 18). In a first approximation, this estimate is consistent with most previous results (Simon et al., 2005; Taylor et al., 2005a, 2005b; Cosarinsky et al., 2006), except those larger than 590,000 years measured in anorthite (Mane et al., 2015) but anorthite is a mineral highly susceptible to Mg isotope re-equilibration and may not provide an accurate age of WLR formation. However, if the range in E101.1 spinel were to correspond to Mg exchange with a reservoir with different ²⁶Al content, the age conclusion would also be erroneous. For instance, a local and variable contribution of interior FUN material devoid of ²⁶Al to the WLR could potentially explain the ²⁶Al variations in WLR spinel. Such an explanation would be in line with the contribution of isotopically heavy Si (and occasionally Mg) from the interior (Fig. 13) and with low $({}^{26}\text{Al}/{}^{27}\text{Al})'_0$ ratios observed in the outer part of profile 1 (Fig. 13c).

We also considered what could be the extent of secondary Al-Mg disturbances in the WLR due to solid state exchange with a reservoir devoid of ²⁶Al. The range of model initial $^{26}Al/^{27}Al$ ratios corresponds to a $\sim 20\%$ exchange for a 1 µm spinel crystal during a 6 months heating at 1000 K (Fig. 18) using the self-diffusion coefficient of Mg in spinel from Sheng et al. (1992). This exchange duration becomes 2.5 My at 750 K and 30 My at 720 K. The composition of the co-existing pyroxene can be further calculated in similar conditions using the self-diffusion coefficient of natural diopside averaged over the crystallographic axes as a first approximation (Zhang et al., 2010) and corresponds to a 11 % exchange for 1

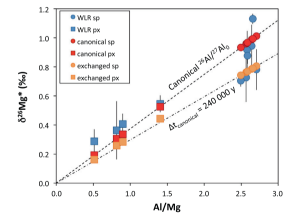


Fig. 18. Comparison between Al/Mg systematics of WLR, theoretical chronology and influence of solid-state diffusion. 2σ errors on measured data (blue). Exchanged values (orange) correspond to solid state diffusion of Mg devoid of ²⁶Mg^{*} during 6 months at 1000 K. Exchanged values are indistinguishable from the isochron expected if crystallization occured 240,000 years after canonical (dash-dotted line). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

µm pyroxene crystals at 1000 K (Fig. 18). At lower temperature and for longer durations, the amount of exchange in the pyroxene becomes closer to that in spinel (e.g., 18% at 720 K). Within analytical error, the pyroxene data cannot be distinguished from a canonical isochron, nor from a younger value, nor from an exchanged value (Fig. 18). This calculation shows that late reheating in the solar nebula (e.g., for the highest temperatures) or parent-body metamorphism (e.g., for the lowest temperatures) are possible sources of chronological disturbance and younger apparent WLR ages because spinel may not have preserved the initial ratio. The range of composition in E101.1 spinel is a possible hint for this process, which may has affected all WLR spinels in other CAIs. We note that 750 K may still be too high for the peak metamorphic temperature undergone by Efremovka for which the maximal temperature is probably closer to ~600 K by analogy with Leoville and Vigarano of petrologic type and thermal history close to those of Efremovka (Huss and Lewis, 1994; Bonal et al., 2006), which would require unreasonably long duration of metamorphism for Mg isotope exchange in spinel. However, interpretation of WLR ages in more metamorphosed meteorites such as Allende must be treated with caution.

Given the uncertainties on the variations of ²⁶Al content in WLR spinel, the initial ²⁶Al/²⁷Al ratio and age of the WLR are probably best given by the converging values of profile 2 and area 4 where the initial value seems best preserved at the contact with the WLR. This initial ratio of $(5.2 \pm 0.1) \times 10^{-5}$ (2 σ) agrees well with that obtained by a regression through WLR data if the three spinel analyses with evidence of disturbances are removed, that is (5.2 \pm $(0.8) \times 10^{-5}$ (2 σ) (Fig. 18). We conclude that the WLR of E101.1 most likely formed with a canonical initial ²⁶Al/²⁷Al ratio. If the canonical ratio corresponds to the time of ²⁶Al homogenization in the early solar system (e.g., Sahijpal et al., 2000; Young et al., 2005; Mishra and Chaussidon, 2014) and the beginning of its chronological use, this constrains the epoch of E101.1 closure to capture of xenolithic materials to within a $\sim 100,000$ years interval. More generally, the measured ages of formation of WLR and their age difference with the interior may not be representative of their true age of formation and many WLR may be essentially contemporaneous with their host CAI formation within the time resolution of the ²⁶Al chronometer. In light of evidence for an isotopic continuum between regular CAIs and FUN CAIs indicated by Ca and Ti isotopes (Park et al., 2014), another important implication of the present work is that mixing with FUN-like precursor material during CAI formation followed by partial melting may be an important source of isotopic disturbance of the Al/ Mg systematics that was previously unrecognized. In this case, many Mg isotopic disturbances, notably in melilite, that were attributed to late secondary processes such as alteration or metamorphism may in fact be of primary nebular origin.

4.7. O isotope reservoirs during E101.1 formation

The coupling of O isotopes with Si and Mg isotope compositions indicates several episodes of ¹⁶O-rich CAI

proto-material dissolution into a ¹⁶O-poor melt. This is clearly the case for (1) the ¹⁶O-rich spinel clusters, which reacted with a ¹⁶O-poor host melt, resulting in local ¹⁶Oenrichments in the gehlenitic products of the reaction and (2) the 16 O-rich sinuous pyroxene fragments, which mixed with a ¹⁶O-poor parental host melt to produce the intermediate compositions observed in areas 3 and 4 and in subinclusion 1. The strong ¹⁶O-depletion observed in the Sc-Zrrich pyroxene, which likely crystallized upon interaction between the host melt and UR perovskite, some of which being ¹⁶O-rich as observed in PerB, further testifies to a strongly ¹⁶O-depleted host melt from which most of the melilite in E101.1 crystallized. The sluggish self-diffusion of O in Ca-rich pyroxene (Ryerson and McKeegan, 1994), requires that this ¹⁶O-depletion is primary. The residual Δ^{17} O heterogeneity within individual areas (Table 2) and the possible oxygen isotope mass fractionation relative to the CCAM line observed in the Sc-Zr-rich pyroxene and some of the ¹⁶O-poor melilite (Fig. 7, Table 2) are well accounted for by assimilation of several ¹⁶O-rich proto-CAIs into a dominantly ¹⁶O-poor melt that had undergone partial evaporation.

All these observations point toward a ¹⁶O-poor CAI precursor at the origin of the E101.1 parental melt (or a strong ¹⁶O-depletion occurring during melting), before the host CAI trapped both the spinel clusters and the sinuous pyroxene xenoliths. The ¹⁶O-depletion must therefore be contemporaneous with the main CAI growth stage by coagulation of proto-CAIs in the solar protoplanetary disk. The Al/Mg systematics of E101.1 further suggests that this growth occurred while the solar system ²⁶Al/²⁷Al ratio was canonical or near canonical. In agreement with increasing evidence (Yurimoto et al., 1998; Aléon et al., 2007; Simon et al., 2011; Kawasaki et al., 2012, 2017; Katayama et al., 2012; Park et al., 2012, 2016; Aléon, 2016), this conclusion indicates that both ¹⁶O-rich (Δ^{17} O $\sim -24\%$) and ¹⁶O-poor ($\Delta^{17}O \sim -2\%$) reservoirs previously evidenced in CAIs (Krot et al., 2002) co-existed in the earliest solar system. The formation of the WLR further constrains the end of E101.1 formation to within the first 100,000 years. At this late stage E101.1 had already experienced a ¹⁶O-depletion, assimilation of ¹⁶O-rich materials and had returned to the ¹⁶O-rich environment, where the AR formed as in many CAIs (e.g., Krot et al., 2002; Yoshitake et al., 2005) and where pyroxene from Type B CAIs crystallized (Aléon, 2016). As shown for several CAIs and expected in many CAIs due to similar general O isotope systematics, the ¹⁶O-depletion was most likely associated with partial melting/crystallization of melilite (Yurimoto et al., 1998; Aléon, 2016; Kawasaki et al., 2017).

Although these observations may be used to constrain the timing and location of oxygen isotope variations in the solar protoplanetary disk, they nevertheless do not shed light on the carrier phases of the oxygen isotope change between solar (McKeegan et al., 2011) and planetary, nor on the physico-chemical processes at the origin of this isotopic difference between solar and planetary reservoirs. Two other observations regarding E101.1 are more pertinent in that respect. First, the primary component related to FUN inclusions is inferred to be dominantly ¹⁶O-poor,

which is in line with the proposition that FUN inclusions formed from a primary dust component of interstellar origin depleted in ¹⁶O relative to the solar composition (Krot et al., 2010; Kööp et al., 2016a). However, it is not clear if the ¹⁶O-poor composition of the FUN component of E101.1 is primary or was acquired during melting by isotope exchange with a ¹⁶O-poor gas of near terrestrial composition. We note that the UR perovskite grains present in the host inclusion were probably initially ¹⁶O-rich, in agreement with the ¹⁶O-rich composition of the sinuous pyroxene, suggesting that most UR materials were initially ¹⁶O-rich. The association of the UR perovskite precursor with the FUN component and the associated ¹⁶O-depletion are the most ancient events that occurred in the history of E101.1 and their record is partially obliterated by subsequent assimilations. We refer to this stage as the "cryptic stage" reflecting that it is not possible to be more precise as to where, when and how the ¹⁶O-depletion occurred.

The sinuous pyroxene fragments contain complex inclusions of secondary phases including wollastonite and oxidized FeO-rich silicates inferred to be andradite and hedenbergite from electron probe analyses by El Goresy et al. (2002). The present discussion is based on this characterization. We show that these secondary FeO-rich minerals are ¹⁶O-poor. They are systematically located in the interior of the sinuous pyroxene and they contributed to the FeO content of the quenched melt region. The host inclusion, by contrast, contains numerous metal inclusions, some of which are quite large. Exsolution of silica, phosphates and V-Fe-oxides (Fig. 4d) from the metal grains is evidence of metamorphism, but most metal grains remain unoxidized. Silicates in the vicinity of metal grains are FeOfree. Although it is widely thought that oxidation and FeO-enrichment of CAI minerals occurred on a parent body in most cases (Brearley and Krot, 2013), these observations strongly suggest that the secondary alteration of the sinuous pyroxene in E101.1 occurred in the protoplanetary disk prior to trapping by the reduced host inclusion (El Goresy et al., 2002). Given that small refractory metal nuggets are present in the sinuous pyroxene fragments, this secondary alteration/oxidation was extremely localized. The conditions of formation of these secondary FeO-silicates are unclear and deserve a detailed study. They point toward existence of a high fO_2 reservoir in the region of CAI formation. This reservoir is not expected from thermodynamic equilibrium considerations (Fedkin and Grossman, 2006; Grossman et al., 2008a) and has a near terrestrial oxygen isotopic composition.

Finally, the relationship between O isotopes and chemistry in sinuous pyroxene (Fig. 8), which shows ¹⁶O depletions associated with less refractory pyroxene (although both variations are small), points toward beginning ¹⁶O depletion in the environment of sinuous pyroxene formation during crystallization. Indeed, the self-diffusion of O in solid pyroxene is sluggish and a solid state secondary re-equilibration, for instance during assimilation of the sinuous pyroxene, would not be correlated with chemistry. This suggests that the ¹⁶O-rich environment in the disk was not entirely homogeneous and that ¹⁶O-rich and ¹⁶Opoor reservoirs somehow interacted with each other. In the vertical zoning model inferred by Aléon (2016), this may result from propagation of the surface ¹⁶O-depletion toward the cool midplane. In the horizontal zoning models such as those inferred from self-shielding associated with the transport of isotopically heavy water ice produced in low temperature regions (e.g., Yurimoto and Kuramoto, 2004; Lyons and Young, 2005; Young, 2007), this may correspond to the mixing front, where ¹⁶O-poor water vapor equilibrates with the ¹⁶O-rich ambient gas. Similar small variations in ¹⁶O-rich hibonite CAIs have been found, which may also point to variable interactions between the ¹⁶O-rich and ¹⁶O-poor gas associated with small temperature differences (Kööp et al., 2016a).

4.8. Protoplanetary disk context of E101.1 formation

Our observations have shown that E101.1 is a compound CAI formed from the coagulation of at least three different precursors (Fig. 19). The first precursor is the ultrarefractory CAI, rich in perovskite, that constitutes the main mass of E101.1. The sinuous pyroxene was related to some extent to this initial UR CAI. It was ¹⁶O-rich and had near canonical abundance of ²⁶Al. Perovskite contained ⁴⁸Ca anomalies. Relative abundances of REE are suggestive of a complex suite of condensation events. including fractional condensation of an UR precursor. The second precursor is a F(UN) inclusion with low ²⁶Al content. The exact conditions of association, melting and evaporation of these two precursors remain unclear because subsequent capture and partial assimilation events obscured these early high temperature stages. After coagulation of these two precursors and their subsequent common thermal history, the main E101.1 host still had a very refractory chemical composition and (i) had experienced a ¹⁶O-depletion, (ii) a strong evaporation, (iii) carried ⁴⁸Ca isotope anomalies, (iv) had heterogeneous ²⁶Al content. After this cryptic stage (Fig. 19a), the main E101.1 captured spinel-rich proto-CAIs, possibly of the common fine-grained spinel-rich CAI type and assimilated the sinuous pyroxene-rich UR CAI. In which order these two capture and partial dissolution events occurred is again not completely clear. Isotopic mixing evidence favors capture of the spinel-CAIs first and sinuous pyroxene last (Fig. 19a). Contrary to the cryptic stage, which can only be inferred from considerations on isotopic and chemical properties of the E101.1 main host, these last two stages are recorded in the petrography of E101.1 and can be visually identified (Fig. 19b). Finally, the WLR formed by condensation into a possibly melted and evaporated outer layer of E101.1 and induced a chemical and isotopic zoning of the outer margins immediately inside the WLR.

It has recently been shown that turbulent transport within the innermost 1 astronomical unit (AU) of a thermally zoned protoplanetary disk before the end of protostellar envelope collapse yields highly variable thermal histories for the various precursors (Taillifet et al., 2014). Extraction of CAIs from the hot zone in such an environment can be realized in at most a few 1000 years. As a result, the various precursors of E101.1 may have collided sequentially with each other during such a turbulent

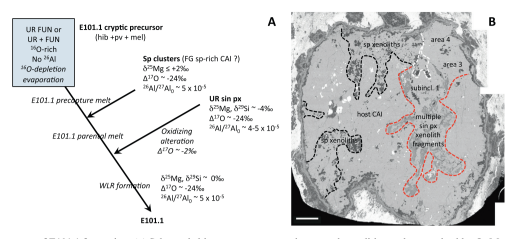


Fig. 19. Main stages of E101.1 formation. (a) Schematic history, precursor and captured xenoliths as characterized by O, Mg and Si isotopes. (b) BSE image of the whole CAI with captured xenolithic material outlined with dashed lines: red - sinuous pyroxene xenoliths, black - spinel clusters xenoliths, white - complex frontier region with possible interactions between sinuous pyroxene, spinel clusters and host. Scale bar 200 μ m. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

journey through the innermost portion of the hot zone of the earliest solar protoplanetary disk. Although not recognized in regular CAIs, the growth of mm to cm sized CAIs through such a series of collisions of numerous precursors may be the standard case of CAI formation since collisions with the nearest neighbors, i.e., between precursors of similar or only marginally different thermal histories, are likely to be statistically favored. The growing identification of compound CAIs supports this conclusion (e.g., El Goresy et al., 2002; Aléon et al., 2005, 2007; MacPherson et al., 2012; Ivanova et al., 2015; Simon et al., 2017). The numerical study of the collisional evolution of CAIs by coagulation and fragmentation in a turbulent hot inner disk indicates rapid growth of mm- to cm-sized objects at collisional equilibrium in a plastic (probably partially molten) state (Charnoz et al., 2015). Such a collisional evolution agrees with aggregation of multiple precursors in E101.1 in a partially molten state and provides a natural explanation for the observed spatial decoupling between different isotope systems and between isotope compositions and mineral chemistry.

Nevertheless, the UR REE and light isotopic composition in the sinuous pyroxene fragments point to a rapid removal of its precursors from the hottest region of the disk and their rapid transport toward the exterior. It remains to be demonstrated to what extent the lack of re-equilibration with the ambient gas fits with the velocities during turbulent transport and the residence time at a given temperature (Taillifet et al., 2014).

Finally, isotopic equilibration between host and xenoliths during capture and assimilation in a partially molten state is a closed-system behavior that requires isolation of the CAI from the ambient gas, notably because the selfdiffusion of O in Ca-Al-rich silicate melts is rapid (Oishi et al., 1974). Mixing relationships are thus expected to be blurred if the partial melt equilibrates with the gas. While this seems not difficult to achieve for Mg and Si due to vastly differing abundances in the CAI and the nebular gas having low partial pressures of Mg and SiO gas, the case of O is more problematic. Physical isolation of melt pockets inside the CAI is unrealistic in light of the distribution of xenolithic fragments and mixed areas, which reach the CAI surface. Chemical isolation is thus required. One way to prevent E101.1 partial melts from equilibrating with the ambient CO or H₂O gas during capture and assimilation (i.e., partial melting) is melting in a diffuse nearvacuum environment with low O abundance and thus low total pressure at solar H/O ratio. Nagahara and Ozawa (2012) calculated that in a closed system with O abundances in the gas about 100 times lower than in a canonical nebula, the final composition of a melt exchanging with the gas is within a few % of the starting composition. However, temperatures in diffuse regions such as the inner edge of the gas disk are likely to be very high and thus to favor a strong evaporation not indicated by the Mg and Si isotopic composition during assimilation of the sinuous pyroxene. In addition, in an open system where the ambient gas can be considered infinite, the partial melt will ultimately equilibrate with the gas, unless the cooling rate is rapid enough to preserve the initial composition. In this case, however, O isotope mass fractionation may be expected depending on the relative timescales of diffusion in the melt and evaporation (Ozawa and Nagahara, 2001). A closed-system behavior of E101.1 during partial melting and partial assimilation of sinuous pyroxene thus remains difficult to explain and to place in an astrophysical context. For that purpose, a better knowledge of isotope exchange rates with relevant astrophysical species such as CO and H₂O in protoplanetary disk conditions (e.g., Yu et al., 1995; Boesenberg et al., 2005, 2016; Di Rocco and Pack (2015)) is required.

5. SUMMARY AND CONCLUSIONS

The multi-isotopic study of E101.1 sheds light on the nature of its various refractory components and on their mixing relationships. Possible precursors have been proposed after disentangling the effects of mixing and partial assimilation. It is found that the final E101.1 CAI results from the coagulation and partial melting of at least three different CAI components:

- 1. An unusual UR CAI dominated by ¹⁶O-rich Al-diopside $(\Delta^{17}O \le -20\%)$ with light Mg and Si isotopic composition (mass fractionation ~-3.5‰/amu in both cases) resulting from kinetic effects, most likely during condensation out of equilibrium. This Al-diopside is now present as multiple relict sinuous pyroxene fragments and locally encloses the ¹⁶O-poor FeO-rich minerals ($\Delta^{17}O$ up to ~-4.5‰) described in El Goresy et al. (2002).
- 2. A melilite-rich UR and FUN host CAI with abundant perovskite, enriched in heavy Mg and Si isotopes by evaporation (δ^{25} Mg and FSi up to $\sim +7\%$ and $\sim +8\%$, respectively). Decoupling of Mg and Si isotopes points toward dissolution of spinel with low δ^{25} Mg after evaporation. Isotopic mixing with the sinuous pyroxene CAI in some regions indicates that the precursor was probably an ultra-refractory melt with little SiO₂ and barely any MgO. The subinclusion 1 identified by El Goresy et al. (2002) is most likely a region of the host that was partially crystallized as gehlenite at the time of sinuous pyroxene capture and remelted subsequently. The calculated pre-evaporation composition indicates that the initial precursor condensed as a hibonite-perovskitemelilite assemblage in agreement with the abundance of melilite and perovskite and evidence of a link with hibonite-rich FUN inclusions.
- 3. The origin of spinel clusters is not as clear but we favor an extraneous source based on several lines of evidence regarding the chemistry, O and Mg isotopes indicating exchange and partial re-equilibration with the host melilite.

The whole CAI is surrounded by a WLR and an accretionary forsterite-rich rim, having isotopic systematics indicative of formation by condensation from a ¹⁶O-rich gas ($\Delta^{17}O \sim -22\%$) with interactions with the evaporated interior. The Al/Mg systematics suggests an early formation (~canonical) of the WLR, approximately contemporaneous of that of the interior CAI with (²⁶Al/²⁷Al)₀ ~ (5.2 ± 0.1) × 10⁻⁵, but also secondary perturbations, which could possibly yield erroneously young WLR formation ages in CAIs if not properly identified ((²⁶Al/²⁷Al)₀ ratios down to 3.7 × 10⁻⁵ in individual spinel grains).

The identification of mixing and interactions between the various precursor CAI components in E101.1 is facilitated owing to the large isotopic differences between the lithological units that result from different thermal histories (e.g., condensation vs evaporation) or from the sampling of different cosmochemical reservoirs (e.g. ¹⁶O-rich vs ¹⁶Opoor or ²⁶Al-rich vs ²⁶Al-poor). Some of these interactions produced isotopic heterogeneities that are also seen in other CAIs with lower amplitude, and remain poorly understood. For instance, many heterogeneities in the Al/Mg systematics are attributed to poorly documented late stage perturbations, but the E101.1 results suggest that they may also result from primary mixing processes between ²⁶Al-rich and ²⁶Al-poor components that are obscured by subsequent partial melting. Compared to the E101.1 precursors that have large isotopic differences, the closer CAI precursors are in chemistry and isotopic composition, the more difficult it is to identify these effects.

The study of E101.1 also sheds light on the origin of O isotopic variations in CAIs. It has been long debated whether CAIs record primary isotopic heterogeneities in the solar protoplanetary disk or underwent secondary perturbations that reset the isotopic composition of some minerals. Recent studies show strong support in favor of the coexistence of ¹⁶O-rich and ¹⁶O-poor reservoirs early in the protoplanetary disk. The O isotope systematics of E101.1 clearly supports these conclusions and demonstrates that igneous CAIs underwent a ¹⁶O-depletion at high temperature contemporaneously with their growth by coagulation.

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APPENDIX A. SUPPLEMENTARY MATERIAL

Supplementary data associated with this article can be found, in the online version, at https://doi.org/10.1016/j. gca.2018.04.001.

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