Refractory inclusions [calcium–aluminum-rich inclusions, (CAIs)] represent the oldest Solar System solids and provide information regarding the formation of the Sun and its protoplanetary disk. CAIs contain evidence of now extinct short-lived radioisotopes (e.g., $^{26}$Al, $^{41}$Ca, and $^{182}$Hf) synthesized in one or multiple stars and added to the protosolar molecular cloud before or during its collapse. Understanding how and when short-lived radioisotopes were added to the Solar System is necessary to assess their validity as chronometers and constrain the birthplace of the Sun. Whereas most CAIs formed with the canonical abundance of $^{26}$Al corresponding to $^{26}$Al/$^{27}$Al of $\sim 5 \times 10^{-3}$, rare CAIs with fractionation and unidentified nuclear isotope effects (FUN CAIs) record nucleosynthetic isotopic heterogeneity and $^{26}$Al/$^{27}$Al of $<5 \times 10^{-3}$, possibly reflecting their formation before canonical CAIs. Thus, FUN CAIs may provide a unique window into the earliest Solar System, including the origin of short-lived radioisotopes. However, their chronology is unknown. Using the $^{182}$Hf–$^{182}$W chronometer, we show that a FUN CAI recording a condensation origin from a solar gas formed coeval with canonical CAIs, but with $^{26}$Al/$^{27}$Al of $\sim 3 \times 10^{-6}$. The decoupling between $^{182}$Hf and $^{26}$Al requires distinct stellar origins: steady-state galactic stellar nucleosynthesis for $^{182}$Hf and late-stage contamination of the protosolar molecular cloud by a massive star(s) for $^{26}$Al. Admixing of stellar-derived $^{26}$Al to the protoplanetary disk occurred during the epoch of CAI formation and, therefore, the $^{26}$Al–$^{26}$Mg systematics of CAIs cannot be used to define their formation interval. In contrast, our results support $^{182}$Hf homogeneity and chronological significance of the $^{182}$Hf–$^{182}$W clock.

Meteorites and their components contain evidence for the presence of now extinct short-lived (t$_{1/2} < 10$ Ma) radioisotopes (e.g., $^{41}$Ca, $^{26}$Al, $^{60}$Fe, $^{53}$Mn, and $^{182}$Hf) during the earliest stages of the Solar System’s evolution. These radioisotopes are believed to have an external, stellar origin, and were either inherited from the ambient interstellar medium or injected into the protosolar molecular cloud before or contemporaneously with its collapse (1). Understanding how and when these radioisotopes were added to the nascent Solar System can constrain the astrophysical environment where our Sun formed and, therefore, test models of Solar System formation. The oldest Solar System solids preserved in chondritic meteorites are calcium–aluminum-rich inclusions (CAIs), which define an absolute age of 4.567 ± 0.16 Ma (2). These millimeter-to-centimeter objects are believed to have formed as fine-grained condensates from a $^{16}$O-rich gas of approximately solar composition in a region with high ambient temperature (>1,300 K) and low total pressures ($\sim 10^{14}$ bar). This environment existed in the innermost part of the protoplanetary disk during the early stage of its evolution characterized by high mass accretion rates ($\sim 10^{-2} M_{\odot}$ y$^{-1}$) to the proto-Sun (3). Formation of CAIs near the proto-Sun is also indicated by the presence in these objects of the short-lived radioisotope $^{10}$Be formed by solar energetic particle irradiation (4). Some of the CAIs subsequently experienced melting and evaporation to form distinct coarser igneous inclusions, such as the compact Type A and Type B CAIs commonly observed in CV meteorites (carbonaceous chondrite of the Vigarano type) (5).

The majority of CAIs in unmetamorphosed chondrites contain high abundance of radiogenic $^{26}$Mg ($^{26}$Mg$^*$), the decay product of $^{26}$Al ($t_{1/2} \sim 0.7$ Ma), corresponding to an inferred initial $^{26}$Al/$^{27}$Al ratio of $\sim (4.5–5.5) \times 10^{-5}$ (6). Recent high-precision $^{26}$Al–$^{26}$Mg systematics of bulk CV CAIs define the so-called canonical $^{26}$Al/$^{27}$Al ratio of $(5.252 \pm 0.019) \times 10^{-5}$ (7). The uncertainty of the canonical $^{26}$Al/$^{27}$Al ratio corresponds to $\sim 4,000$ y, implying a brief episode of condensation and melt evaporation that resulted in Al/Mg fractionation event(s). However, it is uncertain whether the canonical $^{26}$Al/$^{27}$Al ratio reflects that of the bulk Solar System or, alternatively, only a local snapshot of the evolving inhomogeneous protoplanetary disk. A rare subset of refractory grains [platy hibonite crystals (PLACs) and blue aggregates (BAGs), ref. 8] and inclusions (6, 9) have low initial $^{26}$Al/$^{27}$Al ratios ($<5 \times 10^{-5}$). Of particular interest are the coarse-grained igneous inclusions with fractionation and unidentified nuclear effects (FUN CAIs, ref. 10), which, in addition to their low initial abundance of $^{26}$Al, are characterized by large mass-dependent fractionation effects and nucleosynthetic anomalies in several elements. These observations are interpreted to reflect formation of FUN CAIs by thermal processing of presolar dust aggregates before the injection of $^{26}$Al and its homogenization in the protoplanetary disk (11). If this interpretation is correct, FUN CAIs can provide insights into the timing of admixing of $^{26}$Al to the forming protoplanetary disk and, in turn, the origin of short-lived radioisotopes in the early Solar System. However, a late formation of $^{26}$Al-poor CAIs after decay of $^{26}$Al cannot be excluded. Indeed, CAIs are known to have experienced multistage thermal processing in the protoplanetary disk and/or on their chondrite parent bodies (5) that could have erased their radiogenic $^{26}$Mg. Thus, obtaining a robust age estimate for a FUN inclusion is a critical step toward a better understanding of the significance of $^{26}$Al-poor inclusions.
Results and Discussion

Most known FUN CAIs were discovered >30 years ago and were largely consumed during destructive isotopic measurements. To identify additional FUN CAIs suitable for age dating, we conducted a systematic search of coarse-grained refractory inclusions in the Allende CV carbonaceous chondrite, given that CV chondrites contain the highest proportion of igneous CAIs among the distinct chondrite groups. Of ~220 inclusions investigated, only one FUN CAI was identified on the basis of its bulk magnesium–isotope composition. This FUN inclusion, named STP-1, is a coarse-grained igneous Type B2 CAI composed of melilite, spinel, Al,Ti-diopside, and anorthite. STP-1 contains only minor amounts of secondary minerals (nepheline, sodalite, grossular, and monticellite), indicating that it largely avoided secondary alteration processes. Similar to most previously identified FUN CAIs (10, 12), STP-1 shows mass-dependent enrichment in the heaviest isotopes of magnesium, as well as deficits in the mass-independent components of 26Mg (26Mg*) and 54Cr of ~300 and ~3,500 ppm, respectively (Table 1). Trace element analysis demonstrates that STP-1 is characterized by a Group II rare-earth element (REE) pattern (Fig. 1), indicative of condensation from a gas depleted in the most refractory REEs (13). On a three-isotope oxygen diagram, compositions of spinel, anorthite, hibonite, and most Al,Ti-diopside grains plot along a mass-dependent fractionation line with a slope of 0.52 and Δ17O value of ~24 ± 1‰. Oxygen–isotope compositions of melilite and some Al,Ti-diopside grains deviate from this line and show Δ17O values ranging from ~17 to ~4‰ from ~24 to ~17‰, respectively (Fig. 2). The igneous texture and the fractionated magnesium and oxygen–isotope composition favoring the heaviest isotopes imply that STP-1 experienced melt evaporation at low total pressure (14) following condensation of its precursor material. To define the initial abundance of 26Al at the time of crystallization of STP-1, we have investigated the 26Al–27Mg systematics of its primary minerals by secondary ionization mass spectrometry. Multiple analyses of spinel, Al,Ti-diopside, melilite, hibonite, and anorthite crystals define an internal isochron corresponding to an initial 26Al/27Al ratio of (2.94 ± 0.21) × 10−6 (Fig. 3A), that is, much lower than the canonical value of ~5 × 10−5.

Absolute age dating of CAIs by the Pb–Pb method requires knowledge of the uranium–isotope composition of individual inclusions (2). However, the uranium concentration in STP-1 is depleted by a factor of ~100 compared with canonical CAIs, possibly due to loss during melt evaporation under oxidizing conditions. As such, STP-1 in particular, and FUN inclusions in general, may not be suited for uranium-corrected absolute Pb–Pb dating using current state-of-the-art mass spectrometry techniques. To define the formation age of STP-1, we have instead investigated its 182Hf–182W systematics by the internal isochron approach. With a half-life of ~9 Ma, the 182Hf-to-182W decay scheme is one of the most widely used chronometers to understand the timing of solid formation in the early Solar System (15). In addition, currently available data support the proposal that the 182Hf nuclide was uniformly distributed in the early Solar System with an initial 182Hf/180Hf ratio of (9.85 ± 0.40) × 10−5 (16–18). First, the 182Hf–182W isochron of canonical CAIs intersects the carbonaceous chondritic composition, suggesting that the precursor material of primitive asteroids that presumably accreted in the outer Solar System had similar initial 182Hf content as CAIs (16). Second, there is excellent agreement between uranium-corrected Pb–Pb and Hf–W ages of rapidly cooled magmatic meteorites (19). Lastly, CAIs with variable tungsten nucleosynthetic anomalies define the same initial 182Hf abundance (17), indicating that the source of 182Hf is decoupled from that responsible for nucleosynthetic heterogeneity in refractory elements such as tungsten. The Al,Ti-diopside, anorthite and melilite fractions separated from STP-1 define a statistically significant 182Hf–182W isochron corresponding to an initial 182Hf/180Hf ratio of (9.60 ± 1.10) × 10−5 and intercept of ~113 ± 27 ppm, when the 186W/184W is used to correct for instrumental mass fractionation (Fig. 3B). Using a different isotope pair for internal normalization yields an identical initial 182Hf/180Hf within uncertainty but a different intercept of ~26 ± 26 ppm (see the legend of Fig. 3). These two intercept values are distinct from the inferred Solar System initial 182Hf/180Hf value of ~351 ± 10 ppm (18), indicating the presence of tungsten nucleosynthetic heterogeneity in STP-1. In contrast, the internal 182Hf–182W isochron of STP-1 corresponds to a 182Hf/180Hf ratio that is identical within analytical uncertainty to the Solar System initial 182Hf/180Hf ratio of (9.85 ± 0.40) × 10−5 inferred from canonical CAIs (18). Accepting the inferred initial 182Hf/180Hf ratio of STP-1 and associated uncertainty at face value, we calculate an age difference of 0.33 ± 0.67 Ma between formation of STP-1 and canonical CAIs, which is not consistent with the time interval of 3.02 ± 0.07 Ma inferred from the 26Al–27Mg system. Thus, we conclude that the formation age of STP-1 and, by extension, that of 26Al-poor FUN CAIs, is coeval with canonical CAIs within the uncertainty of our measurements. These results are consistent with the proposal that 182Hf was homogeneously distributed in the solar protoplanetary disk at the time of formation of the Solar System’s first solids (16–18).

The contrasting initial abundances of 26Al recorded by CAIs with identical initial 182Hf/180Hf ratios indicate that 26Al was heterogeneously distributed in the protoplanetary disk during the epoch of CAI formation. Therefore, the 26Al–27Mg systematics of CAIs cannot be used to define the duration of the CAI-forming event(s). Likewise, the apparently restricted range of inferred initial 26Al/27Al ratios defined by bulk analyses of canonical CAIs from CV carbonaceous chondrites does not necessarily imply a homogeneous distribution of 26Al throughout the solar protoplanetary disk during and after the epoch of CAI formation. Assessing the degree of 26Al homogeneity in the disk requires careful comparison between the 26Al–27Mg and uranium-corrected Pb–Pb ages of objects with simple thermal histories. We note that the age difference between the formation of canonical CAIs and rapidly cooled angrite meteorites inferred from the assumption-free uranium-corrected Pb–Pb dating method is not consistent with that suggested by the 26Al–27Mg system,

Table 1. 182Hf–182W systematics of mineral fractions and bulk Mg and Cr isotope compositions of the STP-1 FUN CAI

<table>
<thead>
<tr>
<th>Sample</th>
<th>27Al/26Mg</th>
<th>µ26Mg</th>
<th>µ26Mg*</th>
<th>µ53Cr</th>
<th>µ54Cr</th>
<th>182Hf/180W</th>
<th>µ182W (6/3)</th>
<th>µ184W (6/3)</th>
<th>µ182W (6/4)</th>
<th>µ184W (6/4)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk</td>
<td>3.18 ± 0.06</td>
<td>9.331 ± 20</td>
<td>-303 ± 10</td>
<td>260 ± 9</td>
<td>-3.579 ± 15</td>
<td>1.978 ± 0.119</td>
<td>-18 ± 27</td>
<td>-28 ± 15</td>
<td>69 ± 22</td>
<td>41 ± 23</td>
</tr>
<tr>
<td>Melilite</td>
<td>2.675 ± 0.161</td>
<td>35 ± 27</td>
<td>-27 ± 16</td>
<td>115 ± 35</td>
<td>43 ± 24</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anorthite</td>
<td>14.22 ± 0.85</td>
<td>622 ± 52</td>
<td>-31 ± 24</td>
<td>676 ± 58</td>
<td>46 ± 37</td>
<td></td>
<td></td>
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<td></td>
</tr>
</tbody>
</table>

Isotope ratios expressed in the µ-notation, which reflect 106 (ppm) deviations from the terrestrial reference standard. Uncertainties represent the external reproducibility or internal precision, whichever is larger. Mg and Cr isotope data were acquired following techniques outlined in Bizzarro et al. (33) and Trinquier et al. (34), respectively. (6/3), internally normalized to 186W/184W; (6/4), internally normalized to 186W/184W.
supporting widespread $^{26}$Al heterogeneity in the solar protoplanetary disk (2, 7).

The decoupling between the initial abundances of $^{26}$Al and $^{182}$Hf in early-formed refractory inclusions requires distinct stellar sources to account for the presence of these short-lived nuclides in the early Solar System. In addition, the Hf-W data reported here for the STP-1 FUN CAI are most easily understood in the context of a homogeneous distribution of $^{182}$Hf at the birth of the Solar System. If correct, this implies that the carrier of this short-lived radioisotope was well mixed within the Solar System’s parental molecular cloud. Heavy r-process isotopes such as $^{182}$Hf are thought to be synthesized during the explosions of core-collapse supernovae of less than 11 $M_☉$ (20). Because the lifetime of these “lower mass” massive stars is significantly longer than the typical lifetime of giant molecular clouds (21), they are not expected to contribute appreciable amounts of freshly synthesized radionuclides into star-forming regions. Therefore, in agreement with models of the chemical evolution of the galaxy (20, 22), we infer that the initial $^{182}$Hf/$^{180}$Hf ratio of $\sim 1 \times 10^{-4}$ recorded by FUN and canonical CAIs reflects long-term, steady-state galactic stellar nucleosynthesis before the formation of the protosolar molecular cloud. A galactic origin for the initial $^{182}$Hf in the early Solar System is consistent with the view that this radionuclide was homogeneously distributed in the protoplanetary disk at the time of formation of the Solar System’s first solids as inferred from earlier work and the $^{182}$Hf–$^{182}$W data presented here for the STP-1 FUN inclusion.

In contrast with $^{182}$Hf, the Solar System’s initial inventory of $^{26}$Al is approximately 10 times higher than the background levels of the galaxy inferred from γ-ray astronomy (23) and/or models of the galactic chemical evolution (20, 22), requiring late-stage addition of stellar debris to the Solar System’s parental molecular cloud. A possibility is that the observed variable $^{26}$Al abundances during the epoch of CAI formation reflect heterogeneity in the CAI precursor material (24). Such heterogeneity could result from selective thermal processing of presolar carriers, including the carrier(s) of $^{26}$Al, thereby generating reservoirs enriched or depleted in presolar components (25). However, $^{26}$Al-poor objects such as FUN CAIs, PLACs, and BAGs show large-scale nucleosynthetic heterogeneity in the stable $^{48}$Ca and $^{50}$Ti nuclides, including both enrichments and depletions (8–10, 12, 26, 27), implying that the heterogeneity preserved in these objects is unrelated to $^{26}$Al. Moreover, the mineralogy of STP-1, coupled with its group II REE pattern and $^{180}$O-rich composition, suggest that the precursor material of this inclusion formed by condensation from a gas of solar composition depleted in the most refractory REEs, similar to the majority of fine-grained CAIs. Thus, although it is possible that the variable nucleosynthetic anomalies present in $^{26}$Al-poor objects reflect selective thermal processing of their precursors, we conclude that the initial $^{26}$Al/$^{27}$Al of $\sim 3 \times 10^{-4}$ recorded by STP-1 represents the $^{26}$Al abundance in the CAI-forming region when this inclusion crystallized.

Solids ultimately thermally processed in the CAI-forming region are believed to represent molecular cloud material accreting to the proto-Sun from the infalling envelope via the protoplanetary disk (28). In contrast with $^{182}$Hf, our results suggest that the carrier of $^{26}$Al was heterogeneously distributed in the protosolar molecular cloud and, by extension, during infall of envelope material to the protoplanetary disk. However, at the time of formation of the earliest Solar System solids, the innermost protoplanetary disk is thought to have been physically well mixed (29), implying that the observed $^{26}$Al heterogeneity during formation of FUN and canonical CAIs may be temporal rather than spatial. Thus, the different levels of $^{26}$Al in these inclusions could reflect admixing of stellar derived carrier(s) of $^{26}$Al to the protoplanetary disk during the epoch of CAI formation. Progressive admixing of $^{26}$Al to the disk can be understood in the framework of the inside-out collapse model of prestellar cores, where the innermost portion of the core collapses first, followed by the successive outer layers (30). This interpretation requires the innermost part of the protostellar molecular cloud to have been depleted in $^{26}$Al compared with the remaining cloud, and that the formation of FUN CAIs predates canonical CAIs, the latter being allowed by the $^{182}$Hf–$^{182}$W age uncertainty of the STP-1 FUN CAI.

A galactic origin for the initial abundance of $^{182}$Hf recorded by the FUN and canonical CAIs requires $\sim 18$ Ma of free decay between last nucleosynthetic event that produced the heavy
r-process nucleides and formation of the Solar System (22). This time interval, however, is not compatible with the initial \( ^{26}\text{Al} / ^{27}\text{Al} \) ratio of \(-3 \times 10^{-5}\) defined by the STP-1 FUN CAI, accepting that this value represents that of the \(^{26}\text{Al}\)-poor region and the \(^{16}\text{O}\)-rich composition of most primary phases. Thus, we infer that enriched in freshly synthesized matter by earlier generation(s) was part of a giant molecular cloud complex (GMC) chemically.

This interpretation suggests that the protosolar molecular cloud

M.S.W.D., mean square of weighted deviations.

The magnesium from samples with Al/Mg ratios typical of CAIs was

Sample was taken for Al/Mg ratio determination to 5\% accuracy using
diopside fractions were rinsed in distilled ethanol followed by 0.02 M HNO\(_3\) in

Following removal from the Allende slab and cleaning, bulk fragments of

We conducted a systematic search for new FUN CAIs by investigating the Mg–isotope composition of numerous igneous CAI-like objects in cut sections of an \(-3\) kg fragment of the Allende CV carbonaceous chondrite. All igneous CAI-like inclusions of appropriate size were sampled with a computer-assisted

Mixtures of concentrated HF–HNO\(_3\)–H\(_2\)O were loaded on a column containing 2–4 mL of AG50W-X8 resin, and W was eluted along

When the \(^{186}\text{W} / ^{183}\text{W}\) for internal normalization returns an initial \(^{186}\text{Hf} / ^{183}\text{W}\) value of (9.22 \pm 1.1) \times 10^{-5} and an intercept of \(-26 \pm 26\) ppm (M.S.W.D. = 0.31). This \(^{186}\text{Hf} / ^{183}\text{W}\) value is identical within analytical uncertainty to that obtained using the \(^{186}\text{W} / ^{183}\text{W}\) for internal normalization, although marginally lower. However, repeated analysis of a number of distinct aliquots of column-processed BCR-2 (Basalt, Columbia River) rock standard and the Allende carbonaceous chondrite using a quantity of W comparable to that present in the mineral fractions of the STP-1 FUN CAI indicates a superior external reproducibility when the

Following tungsten purification, isotope data were acquired in static mode using the ThermoFisher Neptune multiple collector inductively coupled plasma mass spectrometer (MC-ICPMS) at the Centre for Star and Planet Formation in Copenhagen, following protocols outlined in Bizzarro et al. (33). One inclusion was typified by a resolvable deficit in \(^{26}\text{Mg}\) of \(-300\) ppm as well as a stable Mg–isotope composition enriched in the heavy isotopes by \(-1\%)amu. This inclusion, named STP-1, was classified as a FUN CAI and selected for further analysis. Present on the surfaces of two 3-mm-thick sections, the STP-1 FUN CAI is a spherical inclusion of \(-10\) mm in diameter. Once the inclusion was liberated from the Allende meteorite, polished sections were made from the extracted material for petrographic characterization, mineral chemistry and in situ \(^{26}\text{Al} / ^{27}\text{Al}\)–O-isotope work.

Elemental maps of sections and electron microprobe analyses of individual minerals were performed with the University of Hawaii (UH) field-emission electron JEOL JXA-8500F operated at 15-kV accelerating voltage, 15-nA beam current, and fully focused beam using five wavelength spectrometers. The STP-1 inclusion is a coarse-grained igneous CAI composed of pure anorthite, gehlenitic melilite (\(\text{Åk}_6 \text{Si}_{22}\)), and igneously zoned \(\text{Al},\text{Ti}\)-diopside (\(\text{Al}_9\text{Ti}_9\text{Si}_{29}\text{O}_{55}\)) with 17.7

\(\text{Hf}\)–\(\text{W}\) is used for internal normalization. Therefore, our preferred approach is to use the \(^{186}\text{W} / ^{183}\text{W}\) for internal normalization, in agreement with earlier studies (17, 18, 35, 36).

Fig. 3. Internal mineral \(^{26}\text{Al} / ^{27}\text{Al}\) (A) and \(^{182}\text{Hf} / ^{183}\text{W}\) (B) isochron diagrams for the Allende STP-1 FUN CAI. The well-defined \(^{26}\text{Al} / ^{27}\text{Al}\) isochron based on spinel, \(\text{Al},\text{Ti}\)-diopside, melilite, anorthite, and hibonite shows no evidence for late-stage disturbance, consistent with its pristine mineralogy and the \(^{18}\text{O}\)-rich composition of most primary phases. Thus, we infer that the \(^{26}\text{Al} / ^{27}\text{Al}\) isochron defines the initial \(^{26}\text{Al} / ^{27}\text{Al}\) ratio in STP-1 at the time of its crystallization. M.S.W.D., mean square of weighted deviations.

Using the \(^{186}\text{W} / ^{183}\text{W}\) instead of the \(^{186}\text{Hf} / ^{183}\text{W}\) ratio for internal normalization returns an initial \(^{186}\text{Hf} / ^{183}\text{W}\) value of (9.22 \pm 1.1) \times 10^{-5} and an intercept of \(-26 \pm 26\) ppm (M.S.W.D. = 0.31). This \(^{186}\text{Hf} / ^{183}\text{W}\) value is identical within analytical uncertainty to that obtained using the \(^{186}\text{W} / ^{183}\text{W}\) for internal normalization, although marginally lower. However, repeated analysis of a number of distinct aliquots of column-processed BCR-2 (Basalt, Columbia River) rock standard and the Allende carbonaceous chondrite using a quantity of W comparable to that present in the mineral fractions of the STP-1 FUN CAI indicates a superior external reproducibility when the

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j) were spiked with a mixed \(^{180}\text{Hf} / ^{184}\text{W}\) tracer for elemental abundance determination. The chromium isotope composition of a separate bulk aliquot was determined based on previously published techniques (34) using a ThermoFisher Triton thermal ionization mass spectrometer at the Centre for Star and Planet Formation.

Following handpicking, the anorthite, melilite, and \(\text{Al},\text{Ti}\)-diopside fractions were rinsed in distilled ethanol followed by 0.02 M HNO\(_3\) in

Anorthite, melilite, and \(\text{Al},\text{Ti}\)-diopside fractions were rinsed in distilled ethanol followed by 0.02 M HNO\(_3\) in

No multilayered Wark–Lovring rim sequence is observed around STP-1. The oxygen isotope composition and Al–Mg systematics of primary minerals in STP-1 was investigated using the UH Cameca ims-1280 ion microprobe based on techniques described in SI Materials and Methods.

Following removal from the Allende slab and cleaning, bulk fragments of STP-1 were preserved for bulk isotope and elemental analyses. The remaining material was gently crushed in an agate mortar under distilled ethanol and minerals were handpicked under binocular microscopes in both plain and back lighting. REE abundances were determined on the X-Series II ICPMS from STP-1. The oxygen isotope composition and Al–Mg systematics of primary minerals in STP-1 was investigated using the UH Cameca ims-1280 ion microprobe based on techniques described in SI Materials and Methods.

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The magnesium from samples with Al/Mg ratios typical of CAIs was

Sample was taken for Al/Mg ratio determination to 5\% accuracy using
was applied to all samples, and the uncertainty of this correction is propagated in the final uncertainties of the isotope measurements reported in Table 1. Full analytical details of procedures used for Hf/W and W isotope measurements, as well as all other data reported in this paper, are presented in SI Materials and Methods.