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Rapid condensation of the first Solar System solids

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Chondritic meteorites are composed of primitive components formed during the evolution of the Solar protoplanetary disk. The oldest of these components formed by condensation, yet little is known about their formation mechanism because of secondary heating processes that erased their primordial signature. Amoeboid Olivine Aggregates (AOAs) have never been melted and underwent minimal thermal annealing, implying they might have retained the conditions under which they condensed. We performed a multiisotope (O, Si, Mg) characterization of AOAs to constrain the conditions under which they condensed and the information they bear on the structure and evolution of the Solar protoplanetary disk. High-precision silicon isotopic measurements of 7 AOAs from weakly metamorphosed carbonaceous chondrites show large, mass-dependent, light Si isotope enrichments ($-9\text{‰} < \delta^{30}\text{Si} < -1\text{‰}$). Based on physical modeling of condensation within the protoplanetary disk, we attribute these isotopic compositions to the rapid condensation of AOAs over timescales of days to weeks. The same AOAs show slightly positive $\delta^{25}\text{Mg}$ that suggest that Mg isotopic homogenization occurred during thermal annealing without affecting Si isotopes. Such short condensation times for AOAs are inconsistent with disk transport timescales, indicating that AOAs, and likely other high-temperature condensates, formed during brief localized high-temperature events.

meteorites | condensation | protoplanetary disk | isotopes

The proto-Sun and its surrounding protoplanetary disk formed 4.57 Ga ago from the gravitational collapse of the dense core of a molecular cloud. Chondrites are primitive meteorites left over from the evolution of the Solar protoplanetary disk. They comprise chondrules, fine-grained matrix, and the oldest known solids of the Solar System (1), refractory inclusions in the form of Ca- and Al-rich inclusions (CAIs, millimeter- to centimeter-sized high-temperature complex assemblages of refractory oxides and silicates; ref. 2) and amoeboid olivine aggregates (AOAs, fine-grained aggregates of olivine grains associated with variable proportions of CAI-like materials; ref. 3). The mineralogy of refractory inclusions resembles that of the earliest condensates predicted to form from a cooling gas of solar composition, with AOAs appearing to be somewhat lower-temperature objects than CAIs. This has long upheld the notion that the first solids of the Solar System formed by sequential gas–solid condensation of vaporized presolar matter (4–6), although many have undergone subsequent melting events, such as those which produced the ever-mysterious chondrules (e.g., ref. 7).

However, the timescales of condensation and/or partial evaporation during formation of AOAs and non-igneous CAIs remain largely underconstrained. Mere thermodynamics offer little insight into the “prehistory” of the solids and igneous cooling rates estimates refer to melting/crystallization events (8), not original gas–solid condensation. Al–Mg dating provides only an upper bound of a few 10s or 100s of millennia for the condensation epoch (9–12), which says little on the formation timescales of individual condensates. From an astrophysical viewpoint, evolution of isotherms should be fairly slow (on 10^4 to 10^6 -y timescales; ref. 13) and random motions

between significant condensation fronts should take similarly long (14), so it is often assumed that condensates formed over prolonged timescales (4, 15, 16). In contrast, Sugiura et al. (17) inferred from the absence of low-Ca pyroxene on most AOAs cooling rates >0.02 K/h at the end of olivine condensation and Komatsu et al. (18) derived 50 K/h from the presence of silica in one unusual AOA.

Stable isotopes may provide insights into the kinetics of gas–solid processes (19–21). Lighter isotopes are more gas-mobile and will thus preferentially impinge a solid grain, giving it an isotopically light composition provided that back-evaporation has not erased the initial fractionation. However, CAIs commonly show mass-dependent, heavy Si, Mg, O signatures (22) diagnostic of partial evaporation of refractory liquids resulting from the melting of solar condensates (23)—whose formation conditions are hereby blurred.

Yet the relatively fine-grained and sometimes porous AOAs have never been melted and have undergone only moderate thermal annealing during the evolution of the Solar protoplanetary disk (3, 24), implying they might represent fairly pristine condensates. Moreover, because AOAs record the condensation of olivine, which represented 1/3 of CI (Ivuna-like) chondritic matter (25)—second only to ice in terms of sheer mass—they are more representative of the protoplanetary disk than CAIs. However, in situ stable isotopic characterizations of AOAs have been performed only for oxygen, an element notoriously subject to poorly understood mass-independent fractionations within the protoplanetary disk, potentially blurring any mass-dependent condensation signal (26, 27). Here, we report the oxygen, silicon, and magnesium isotopic composition of Mg-rich olivine grains in AOAs from a set of carbonaceous chondrites characterized by minimal thermal metamorphism. We use our results to quantify the conditions under which primordial dust formed in the Solar System and discuss the

Significance

Combined O, Si, Mg isotopic measurements of amoeboid olivine aggregates allow the individual condensation timescales of a typical set of the first Solar System solids to be estimated. Our results indicate formation over days or weeks, much faster than could be inferred from the secular evolution of the Solar protoplanetary disk. The oldest solids of the Solar System thus bear witness to a turbulent disk with strong thermal heterogeneities.

Author contributions: Y.M. and J.V. designed the study and performed the isotopic measurements; Y.M., J.V., E.J., M.P., and M.C. discussed the ion probe data; E.J. devised the astrophysical model; Y.M., J.V., E.J., and M.C. wrote the paper.

The authors declare no competing interest.

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Data deposition: All raw SIMS data, summary of the data, and output of the model can be found at <https://data.mendeley.com/datasets/dnxgdbzmzk/3>.

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implications on the structure and evolution of the Solar protoplanetary disk.

Material and Methods

We surveyed all AOAs in a section each of Kaba (thin section N4075 from the Natural History Museum, Vienna, Austria), Northwest Africa (NWA) 5958 (thick section NWA 5958-1 from the Muséum national d'Histoire naturelle, Paris, France) and Miller Range (MIL) 07342 (thick section MIL 07342,9 from the NASA Antarctic Search for Meteorites program). Kaba is an oxidized Bali-like CV chondrite (28, 29). MIL 07342 is a CO chondrite. Although opaque assemblages are altered and evidence of incipient metasomatism appears on chondrule mesostasis, olivine compositions in type I chondrules (e.g., $\text{Fo}_{94.3-99.5}$) show no diffusional exchange; the chondrite is deemed of type 3.0–3.2 by the Antarctic Meteorite Petrographic Description database. NWA 5958 is a C2-ung CM-like chondrite, whose type II chondrule olivine Cr content as well as opaque petrography indicate minimal thermal metamorphism (<300 °C), yet chondrule mesostases have undergone extensive aqueous alteration, which did not affect olivine (30, 31).

Scanning electron microscope observations of AOAs were performed at CRPG-CNRS (Nancy, France) using a JEOL JSM-6510 with 3-nA primary beam at 15 kV. Quantitative compositional analyses of olivine in AOAs were performed using a CAMECA SX-100 electron microprobe [at CAMPARIS (Sorbonne Université, Paris, France)]. A 20-nA focused beam accelerated to 15-kV potential difference was used for spot analyses of olivine with 20-s analysis times. The PAP software was used for matrix corrections.

We analyzed the silicon isotopic compositions of olivines by secondary ion mass spectrometry (SIMS) using the multicollector CAMECA IMS 1270 E7 at CRPG. Olivines were sputtered with an ~5-nA primary Cs^+ beam set in Gaussian mode and accelerated at 10 kV (32). Secondary negative (28–30) Si^- ions were accelerated at 10 kV and analyzed in multicollection mode on 3 off-axis Faraday cups (L'2, C, and H1, respectively). Charge accumulation on the sample surface was neutralized with an electron gun. The mass-resolving power was set at $M/\Delta M = 5,000$ (slit 2) to completely resolve interferences on masses, refs. 28–30. The yields and backgrounds of the Faraday cups were calibrated at the beginning of each analytical session. Automatic centering of the transfer deflectors and mass was implemented in the analysis routine. A $10 \times 10\text{-}\mu\text{m}^2$ raster was applied to the primary beam to ensure flat-bottomed pits. We used 4 terrestrial standard materials (San Carlos olivine, synthetic forsterite, quartz, diopside) to define the terrestrial fractionation line. As there is no matrix effect for Si isotopes for olivine with $\text{Mg\#}$ ($= 100 \times \text{Mg}/[\text{Mg} + \text{Fe}]$) above 70 (32), we used the San Carlos olivine to correct for instrumental mass fractionation. Measurements typically consisted of a 90-s pre-sputtering, automatic mass and beam centering, and 50 cycles of 4-s integrations separated by 1-s

waiting times. Thus, each measurement took ~7 min. Under these conditions, internal precision on $\delta^{29}\text{Si}$ and $\delta^{30}\text{Si}$ was ± 0.05 through 0.30‰ and ± 0.10 through 0.60‰ (2σ SE), respectively, depending on the sample, and the external reproducibility on $\delta^{29}\text{Si}$ and $\delta^{30}\text{Si}$ for the San Carlos olivine was $\pm 0.10\text{‰}$ and $\pm 0.15\text{‰}$, respectively (2σ SE). We also analyzed the oxygen and magnesium isotopic compositions of olivines in the same AOAs following previously established protocols (see *SI Appendix* for technical details).

Results

AOAs in NWA 5958 and MIL 07342 have compact textures characterized by continuous olivine (Fig. 1 and *SI Appendix*, Figs. S1 and S2) (33). Ca–Al-rich vermicular patches are altered, although diopside persists, in AOAs in NWA 5958. AOAs in Kaba show smaller olivine zones, diopside, and Ca–Al-rich areas (*SI Appendix*, Fig. S3). Olivine in all AOAs investigated herein are Mg-rich with $\text{Mg\#}$ in the range 98.9 to 99.9 (*SI Appendix*, Table S1). AOA olivine grains have O isotopic compositions with $\delta^{18}\text{O}$ values ranging from -48.6 to -39.8‰ , $\delta^{17}\text{O}$ values ranging from -47.7 to -41.5‰ , and $\Delta^{17}\text{O}$ values ranging from -23.9 to -19.2‰ (*SI Appendix*, Fig. S4 and Table S2). The silicon isotopic compositions of AOA olivines show large, mass-dependent isotopic variations with $\delta^{29}\text{Si}$ values ranging from -5.1 to -1.0‰ and $\delta^{30}\text{Si}$ values ranging from -9.7 to -0.2‰ (Figs. 1 and 2 and *SI Appendix*, Table S3). We did not observe any correlation between the Si compositions and the analytical location within the AOAs (Fig. 1 and *SI Appendix*, Figs. S1 and S2). The magnesium isotopic compositions of olivines in AOAs show slightly positive $\delta^{25}\text{Mg}$ values ranging from 0.09 to 1.81‰ (Fig. 3 and *SI Appendix*, Table S4). We did not observe any correlation between the average Si and Mg isotopic compositions of the AOAs, nor with their elemental compositions.

Duration and Contexts of AOA Condensation. The mass-dependent character of the Si isotopic variations precludes their explanation by nucleosynthetic anomalies in the AOA-forming region. Provided bulk carbonaceous chondrites are a suitable proxy for the composition of the reservoir from which AOAs formed, the light isotope enrichment is also at variance with the positive $2\text{‰}/u$ fractionation expected from olivine-gas equilibrium (34) and inferred by ref. 15 to account for the Si isotopic variations of bulk chondrite. Partial evaporation would likewise produce heavy compositions in olivine, unless it occurred subordinately

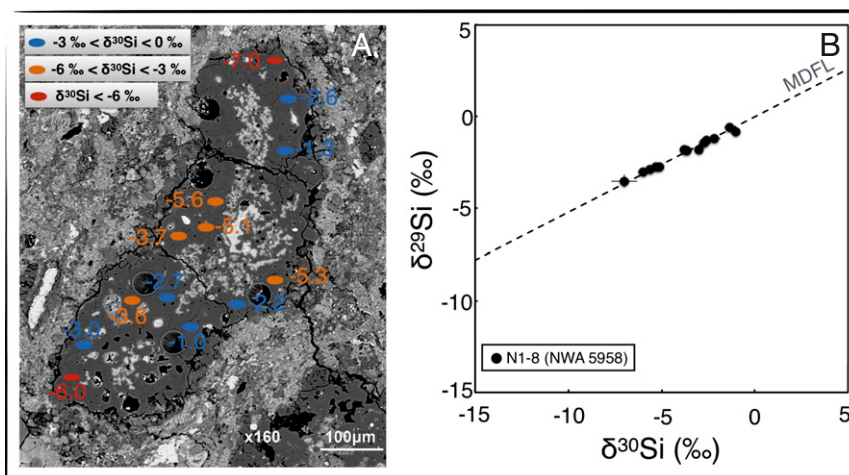


Fig. 1. A representative AOA and the Si isotopic compositions of its olivines. (A) Back-scattered electron image of AOA N1-8 from the CM-related chondrite NWA 5958 (30, 31). SIMS measurements are indicated by colored ellipses and the Si isotopic compositions associated with each analysis point. The color of the ellipses indicates the variability of the Si isotopic compositions by the range of the $\delta^{30}\text{Si}$ value: blue, $-3\text{‰} < \delta^{30}\text{Si} < 0\text{‰}$, orange, $-6\text{‰} < \delta^{30}\text{Si} < -3\text{‰}$, and red, $\delta^{30}\text{Si} < -6\text{‰}$. (B) $\delta^{29}\text{Si}$ – $\delta^{30}\text{Si}$ diagram showing the mass-dependent variation observed in N1-8; MDFL, mass-dependent fractionation line.

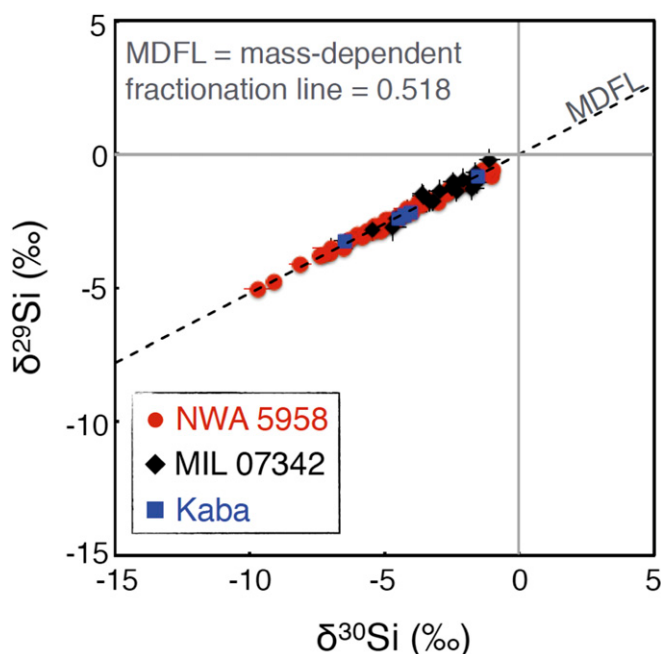


Fig. 2. Silicon 3-isotope plot showing the large range of mass-dependent isotopically light values measured in all analyzed AOAs in NWA 5958 (red), MIL 07342 (black), and Kaba (blue).

on isotopically lighter objects. In contrast, light isotopic enrichment would be expected from condensation, as the light isotopomers of gaseous SiO would collide with the growing solids at greater rates proportional to their thermal velocity (e.g., ref. 20)

$$v_{T, \text{SiO}} \equiv \sqrt{\frac{8kT}{\pi m_{\text{SiO}}}},$$

where k is the Boltzmann constant, T is the temperature, and m_{SiO} is molecular mass. We note that the greatest observed Si isotopic fractionations, around -5‰ /u (Fig. 2 and *SI Appendix, Table S3*), are not far from the maximum kinetic fractionations of -11‰ /u, assuming that the condensation coefficient γ_{SiO} has no mass dependence (*SI Appendix, Fig. S5*). This indicates back-evaporation has had little opportunity to curtail the isotopic fractionation of the condensate. This is possible if the reservoir temperature evolved through the olivine condensation interval over a time shorter than the “SiO condensation timescale,” that is, the theoretical (e-folding) time of SiO condensation (on the available solids) in the absence of evaporation, given by (20) (*SI Appendix*):

$$t_{\text{cond}} \equiv \frac{1}{n_p \pi a^2 \gamma_{\text{SiO}} v_{T, \text{SiO}}},$$

where n_p is the number density of condensate particles, assumed to be identical spheres of density $\rho_s = 3 \times 10^3 \text{ kg/m}^3$ and radius a . If indeed the reservoir temperature evolved more rapidly than t_{cond} , high supersaturation would develop and make back-evaporation subdominant. In Fig. 4, we show the distribution of $\delta^{30}\text{Si}$ expected for different cooling timescales λ^{-1} (λ being defined as the decay constant of the saturation density of Mg, related by stoichiometry to SiO; see *SI Appendix*). It is seen that only $\lambda t_{\text{cond}} > 1$, that is, λ^{-1} values shorter than t_{cond} , can reproduce the range of the data. Thus, AOAs must have formed over a timescale of $\max(t_{\text{cond}}, \lambda^{-1}) = t_{\text{cond}}$.

So what is the magnitude of t_{cond} ? For a steady α disk of mass accretion rate $\dot{M}$, using the pressure expression (*SI Appendix*), we have

$$\begin{aligned} t_{\text{cond}} &= \frac{m_{\text{SiO}}^{1/2}}{32 \gamma_{\text{SiO}} m_g^{5/6} T^{7/3}} \left(\frac{k \rho_s a \dot{M}}{\epsilon_p} \right)^{1/3} \left(\frac{9\pi}{\sigma} \right)^{2/3} \\ &= 10^5 \text{ s} \left(\frac{0.1}{\gamma_{\text{SiO}}} \right) \left(\frac{a}{30 \mu\text{m}} \right)^{1/3} \left(\frac{\alpha}{10^{-2}} \right)^{1/3} \left(\frac{\dot{M}}{10^{-7} M_\odot/\text{a}} \right)^{1/3} \\ &\quad \times \left(\frac{10^{-3}}{\epsilon_p} \right)^{1/3} \left(\frac{1400 \text{ K}}{T} \right)^{7/3}, \end{aligned}$$

where α is the turbulence parameter (35), σ is the Stefan–Boltzmann constant, $m_g = 2.34 \text{ u}$ is the mean nebular gas molecular mass, ϵ_p is the solid/gas mass ratio, and $M_\odot$ is the solar mass. Quite insensitively to the values of the disk parameters, our calculations thus reveal a short duration on the order of a day ($\sim 10^5 \text{ s}$) for the condensation of AOAs, or weeks if smaller values of γ_{SiO} apply (e.g., ref. 36 and Fig. 4). This is consistent with the cooling rate lower limit of 0.02 K/h derived by (17) from the quasisystematic absence of low-Ca-pyroxene in AOAs, as well as the 50 K/h cooling rate suggested for the silica-bearing AOA of ref. 18. This may also account for the disequilibrium gas–solid condensation effects described for AOAs in Allan Hills 77037 (24).

Such durations are, however, much shorter than the 10^4 to 10^6 -y timescales of secular evolution (and cooling) of the disk as a result of progressive accretion into the Sun (13). They are also much shorter than radial transport timescales, which ref. 37 estimated on the order of 10^3 y (*SI Appendix*). Even vertical transport toward cooler surface layers, away from the disk mid-plane, would take much longer than the orbital period (38), even if winds are taken into account (*SI Appendix*). However, the smooth temperature profiles generally invoked in the inner disk are only average; heating would have been dictated by the dissipation of turbulence and likely prone to local heterogeneities, for instance in current sheets in magnetorotational instability-driven turbulence (39–41). Outside the “nominal” condensation front of forsterite, transient temperature excursions may have led to the olivine vaporization and recondensation. Perhaps late mixing with isotopically unfractionated SiO from the surroundings prevented the positive isotopic compositions predicted at the end of closed-system condensation (Fig. 4) but not observed in the data, although we cannot exclude an effect of the finite size of the ion probe beam (*SI Appendix*). In our model (illustrated schematically in Fig. 5), AOAs would have formed at

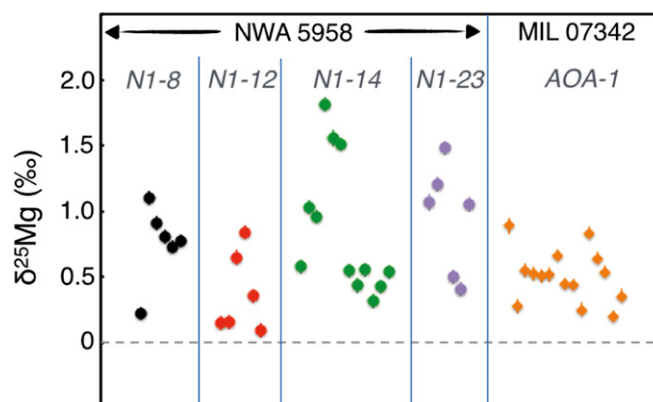


Fig. 3. $\delta^{25}\text{Mg}$ values of AOAs analyzed in NWA 5958, MIL 07342, and Kaba.

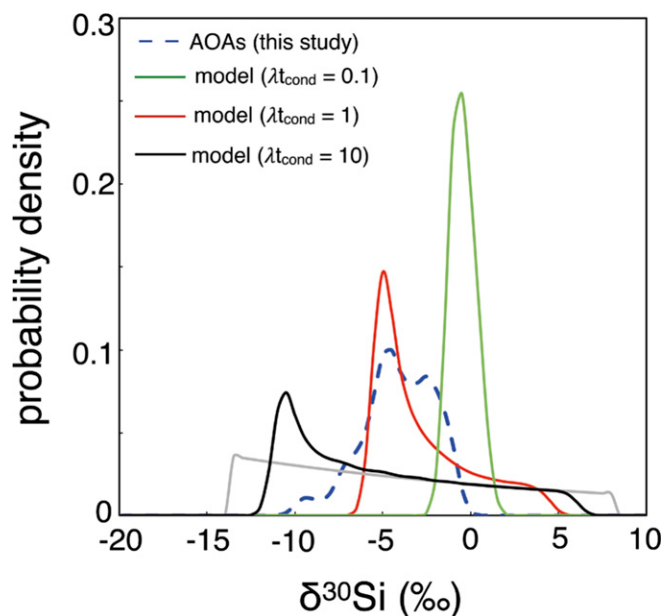


Fig. 4. Probability density of the $\delta^{30}\text{Si}$ values of the studied AOA (blue) compared to a closed-system condensation model. The model curves correspond to different speeds of evolution of the system, measured by the dimensionless parameter $\lambda t_{\text{cond}} = -t_{\text{cond}} d \ln n_{\text{sat}} / dt$ (where n_{sat} is the saturation concentration of Mg; *SI Appendix*), with values of $\lambda t_{\text{cond}} = 0.1$ (green), 1 (blue), 10 (black), and ∞ (gray). The range of Si isotopic fractionations observed herein can only be reproduced when $\lambda t_{\text{cond}} > 1$, although different thermal histories are likely convolved in the measured distribution.

greater heliocentric distances than a mere consideration of the average isotherms would suggest. Those which we now see in chondrites would have eventually escaped the evaporation/recondensation cycle of their matter during outward migration toward less-heated regions, e.g., as a result of turbulent diffusion or disk expansion (13, 42, 43). The “frozen-in” AOA would then have eventually reached the cold regions of chondrite accretion.

Similar processes would likely hold for higher-temperature condensates, i.e., CAIs, at shorter heliocentric distances. There may thus be a continuum between the condensation processes and subsequent reheating events that led to their remelting and crystallization, with cooling rates on the order of 0.1 to 10 K/h (8)—not to mention chondrule-forming processes as well, with cooling rates estimates spanning 1 to 3,000 K/h (ref. 44 and references therein)—the formation of Wark–Lovering rims (45–47) and/or partial evaporation resulting in heavy isotopic enrichments (19). The isolation of the ultrarefractory component presumably lost from the parent reservoir of numerous group II CAIs (48) may have likewise required similarly short timescales. Indeed, the light isotopic enrichment of (refractory) heavy rare-earth elements in such CAIs (49) allows for only limited recondensation during the partial evaporation of the now-lost ultrarefractory complement.

Condensate Agglomeration. It is instructive to compare the condensation timescale calculated above to the grain–grain collision timescale t_{coll} —that is, the mean time between 2 collisions for a given grain, given by ref. 50:

$$t_{\text{coll}} \equiv \frac{1}{n_p 4\pi a^2 \Delta v_{\text{coll}}},$$

where Δv_{coll} is the typical particle–particle velocity. Comparing the 2 timescales, we obtain

$$\frac{t_{\text{coll}}}{t_{\text{cond}}} = \frac{\gamma_{\text{SiO}} v_{T,\text{SiO}}}{4 \Delta v_{\text{coll}}} = 20 \left(\frac{\gamma_{\text{SiO}}}{0.1} \right) \left(\frac{1 \text{ m/s}}{\Delta v_{\text{coll}}} \right) \left(\frac{T}{1,400 \text{ K}} \right)^{1/2}.$$

Since Δv_{coll} is normalized to an upper bound (the critical fragmentation or bouncing velocity; ref. 51), t_{coll} is resolvable longer than t_{cond} , implying that aggregation of individual condensates, if colliding below the critical velocity for agglomeration [which might have been greater for higher temperatures (52)], generally occurred after, not during, condensation (Fig. 5). Indeed, unmelted fine-grained CAIs are composed of mineralogically zoned sub-100- μm nodules with more refractory minerals concentrated in their center, consistent with an origin as individual condensates, which encompassed the whole refractory condensation sequence before coagulation (53). Still, because t_{coll} should be much shorter than radial or vertical transport timescales (14), agglomeration should have occurred largely between genetically related bodies. This accounts for the generally fairly uniform character of fine-grained CAIs at mesoscale, notwithstanding common macroscale layering (53, 54). Same holds for AOA. One can envision that the high-temperature events that ultimately led to their final condensation likely first evaporated all but the most refractory (CAI-like) phases of any dust grains present. Olivine recondensation would have then mantled these dust grains; perhaps some olivine also condensed homogeneously (such as the ^{16}O -rich ones found in chondrite matrices, refs. 55 and 56). These different olivine-bearing bodies would have then aggregated together. While, as argued above, aggregated bodies might have genetic relationships to one another, they may represent different thermal events, if t_{coll} was longer than transport timescales or evaporation/recondensation cycles, and thus possibly variable thermal histories. Thus, individual AOA may group bodies with similar or various ranges in Si isotopic fractionation. Depending on the recondensation events sampled by individual AOA, different AOA may vary in their ranges of $\delta^{30}\text{Si}$.

Secondary Heating of Amoeboid Olivine Aggregates. Since $\delta^{25}\text{Mg}$ corresponds to similar isotopomer mass ratios as $\delta^{30}\text{Si}$ (in terms of Mg and SiO, respectively), one would expect similar fractionations for the former in AOA (*SI Appendix*). Mg isotope fractionations as large as -14‰ have been reported in fine-grained CAIs in Allende (57, 58). However, we observe little variation of $\delta^{25}\text{Mg}$ in the AOA analyzed herein (only 0.4‰ SD), although (59) reported $\delta^{25}\text{Mg}$ down to -3‰ in some porous AOA in Kaba. The compact textures of our AOA suggest significant postagglomeration annealing, which may have led to the Mg isotopic homogenization given the much higher diffusivity of Mg relative to Si (60). Indeed, for a Mg diffusion coefficient of $\sim 10^{-17} \text{ m}^2/\text{s}$ at 1,400 K, isotopic homogenization over lengths of 50 μm requires only an integrated (possibly discontinuous) time of a decade or so (Fig. 5), whereas O and Si homogenization requires durations of 6 orders of magnitude longer. A timescale of a decade would agree with 1) the $<60 \text{ y}$ of isotopic exchange with ^{16}O -poor gas for 2 Leoville AOA (61) and 2) the effective cooling rate of 0.002 K/h estimated from the closure of Mn and Cr diffusion in AOA olivine (17). Ca from the Ca–Al-rich nodules would have diffused as well into forsterite (17, 31), accounting for its apparent Ca enrichment in compact AOA (3, 17, 31), which our Si isotopic results no longer allow to explain by a condensation slower than for porous AOA (17).

The closure of the Mg isotopic system in AOA would post-date the condensation and aggregation period but would not have been as late as parent body metamorphism, because AOA retain near-canonical Al–Mg mineral isochrons (e.g., refs. 62 and 63). The Mg isotopic composition of the bulk olivine (to which all olivines would have been homogenized) would have been also closer to the reservoir average than that of Si because Mg would be essentially completely condensed during olivine condensation,

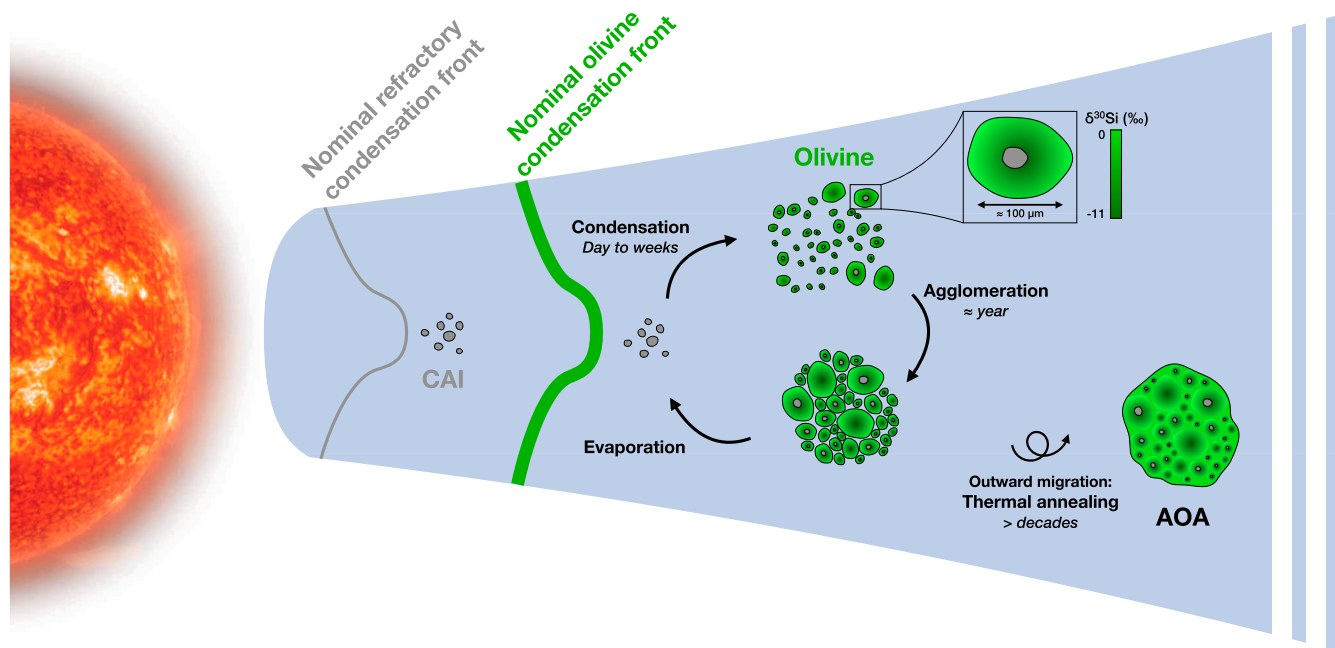


Fig. 5. Schematic representation of AOA formation. Localized heating beyond the nominal olivine condensation front may have evaporated solids, leaving refractory residues. Upon cooling, such CAI-like materials served as seeds for olivine condensation slightly beyond the nominal olivine condensation front. The rapid condensation inferred herein (day to weeks) would have produced large negative mass-dependent silicon isotopic fractionations. Agglomeration of AOAs occurred over a longer timescale on the order of a year. Subsequent thermal annealing produced the AOAs as we observe them today, homogenizing their Mg isotopic compositions without affecting their silicon isotopic compositions.

while almost half of the original SiO would still remain (ref. 6 and *SI Appendix*).

In conclusion, our results suggest that the earliest solids of the Solar System condensed, aggregated, and were processed over short timescales, perhaps on the order of days, in a dynamic and thermally nonuniform disk. Nowhere was the primordial condensation sequence quiescent.

Data Availability. All of the data are available in the *SI Appendix* (*SI Appendix*, Tables S1–S4). All raw SIMS data, summary of the data, and output of the model can be accessed here: <https://data.mendeley.com/datasets/dnxgdbzmzk/3>.

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