



# A 4,565-My-old andesite from an extinct chondritic protoplanet

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**The age of iron meteorites implies that accretion of protoplanets began during the first millions of years of the solar system. Due to the heat generated by <sup>26</sup>Al decay, many early protoplanets were fully differentiated with an igneous crust produced during the cooling of a magma ocean and the segregation at depth of a metallic core. The formation and nature of the primordial crust generated during the early stages of melting is poorly understood, due in part to the scarcity of available samples. The newly discovered meteorite Erg Chech 002 (EC 002) originates from one such primitive igneous crust and has an andesite bulk composition. It derives from the partial melting of a noncarbonaceous chondritic reservoir, with no depletion in alkalis relative to the Sun's photosphere and at a high degree of melting of around 25%. Moreover, EC 002 is, to date, the oldest known piece of an igneous crust with a <sup>26</sup>Al-<sup>26</sup>Mg crystallization age of 4,565.0 million years (My). Partial melting took place at 1,220 °C up to several hundred kyr before, implying an accretion of the EC 002 parent body ca. 4,566 My ago. Protoplanets covered by andesitic crusts were probably frequent. However, no asteroid shares the spectral features of EC 002, indicating that almost all of these bodies have disappeared, either because they went on to form the building blocks of larger bodies or planets or were simply destroyed.**

andesitic magmatism | early solar system | planetary differentiation | differentiated meteorites

Despite the large number of samples in the meteorite record that originate from the crust or mantle of rocky bodies (about 3,100 are known today), these rocks provide an incomplete picture of the diversity of the differentiated bodies that formed in the early solar system (1). Indeed, about 95% of these meteorites originate from only two bodies, with 75% coming from the crust of a single asteroid (possibly 4 Vesta) and the other 20% from the mantle of a presumably larger object, the now-destroyed ureilite parent body (2, 3). Thus, until recently, known achondritic lavas were essentially basalts (eucrites) from 4-Vesta and a handful of other basaltic rocks from unknown parent bodies [the angrites and some ungrouped achondrites such as Northwest Africa (NWA) 011 (4) or Ibitira (5)]. Although certainly not representative of the magmatic activity of all the planetesimals, these achondritic lavas strengthened the general view that their crusts were essentially basaltic in composition. However, the discovery of some rare achondrites of andesitic or trachyandesitic composition [e.g., Graves Nunataks 06128 and 016129 (6, 7), ALM-A (8), NWA 11119 (9)] demonstrated that the diversity of the lavas formed on protoplanets may have been more important than previously thought. Experimental studies motivated by these new meteorites have shown that the generation of silica-rich liquids is possible from the melting of chondrites (10–12). Thus, the formation of andesitic crust was possibly common on protoplanets, especially for those that were not Na and K depleted (12), contrary to what the meteorite record suggests. However, the processes that built such a crust, and the genesis of protoplanetary andesites are not well known due to the rarity of the samples. Here, we report on

Erg Chech 002 (EC 002), a unique andesite achondrite found in the spring of 2020 in the Sahara. This meteorite is the oldest magmatic rock analyzed to date and sheds light on the formation of the primordial crusts that covered the oldest protoplanets.

## Results

**Petrography.** EC 002 is an unbrecciated greenish rock with a medium-grained groundmass (grain size in the order of 1 to 1.5 mm) with 8 vol% pores. Its texture suggests a crystallization within a thick flow or in a shallow intrusion (Fig. 1). It consists of 45 vol% of lath-shaped albitic feldspar (plagioclase and possibly anorthoclase) containing lamellae of K-feldspar, 38 vol% of anhedral pyroxene (ca. 0.6 × 3.7 mm), 5 vol% interstitial silica minerals (cristobalite and tridymite [PO] identified by Raman spectroscopy [*SI Appendix, Figs. S1 and S2*] with no quartz), and minor spinel, ilmenite, Ca-phosphate, troilite, and FeNi metal. The samples we have examined are rather fresh, with only a few rust patches produced by the alteration of troilite and metal and some carbonate fillings in the fractures, which are typical of Saharan finds (13). Pyroxenes are partly equilibrated and consist of relict augite with fine, closely spaced (~1 μm thick) exsolution lamellae of low-Ca pyroxene, or relict low-Ca pyroxene with fine exsolution lamellae of augite. In addition, EC 002 also contains angular to rounded pyroxene and olivine xenocrysts, irregularly dispersed in the groundmass (Figs. 1 and 2) and reaching several centimeters [up to 9 cm (14)]. They are generally mantled by fine-grained groundmass pyroxenes. EC 002 is slightly shocked. Feldspar and pyroxene show only moderate mottled extinction. Some feldspar grains have fine polysynthetic twinning. Fractures in these minerals are not prominent. The shock stage is M-S2 (15).

## Significance

The crusts of the oldest protoplanets are virtually unknown due to the scarcity of samples. Here, we describe the oldest known lava that crystallized ca. 4,565 Ma ago and formed by partial melting of a chondritic parent body. <sup>26</sup>Al-<sup>26</sup>Mg systematics suggest that the elapsed time between melting and crystallization was significant, on the order of several 10<sup>5</sup> y, probably due to the viscosity of the magma. Although the first protoplanetary crusts were frequently not basaltic, their remains are not detected in the asteroid belt because their parent bodies served as the building blocks for larger rocky bodies or were nearly totally destroyed.

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The authors declare no competing interest.

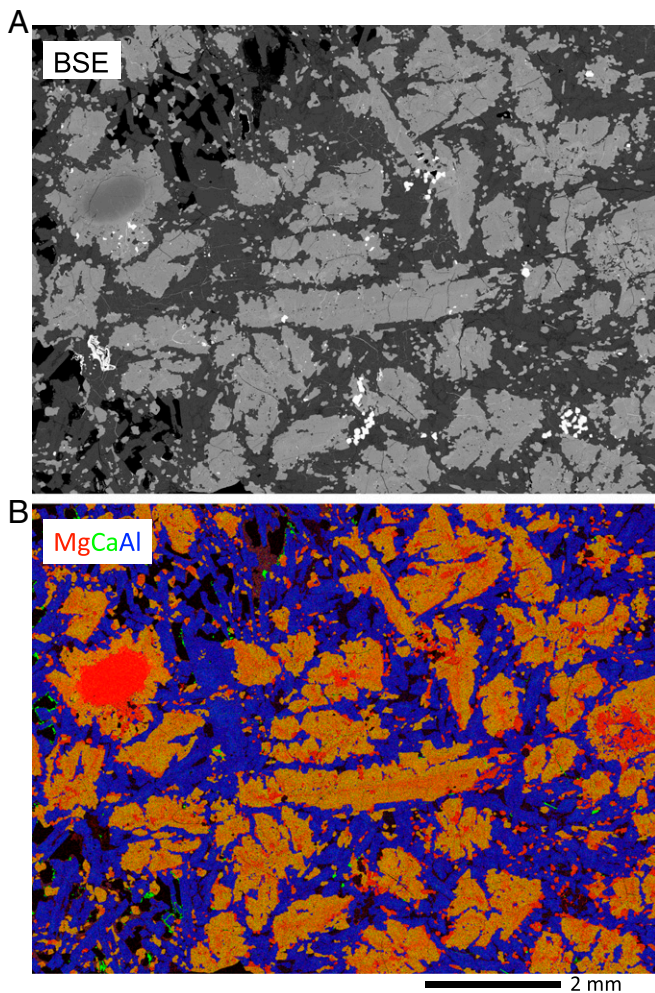
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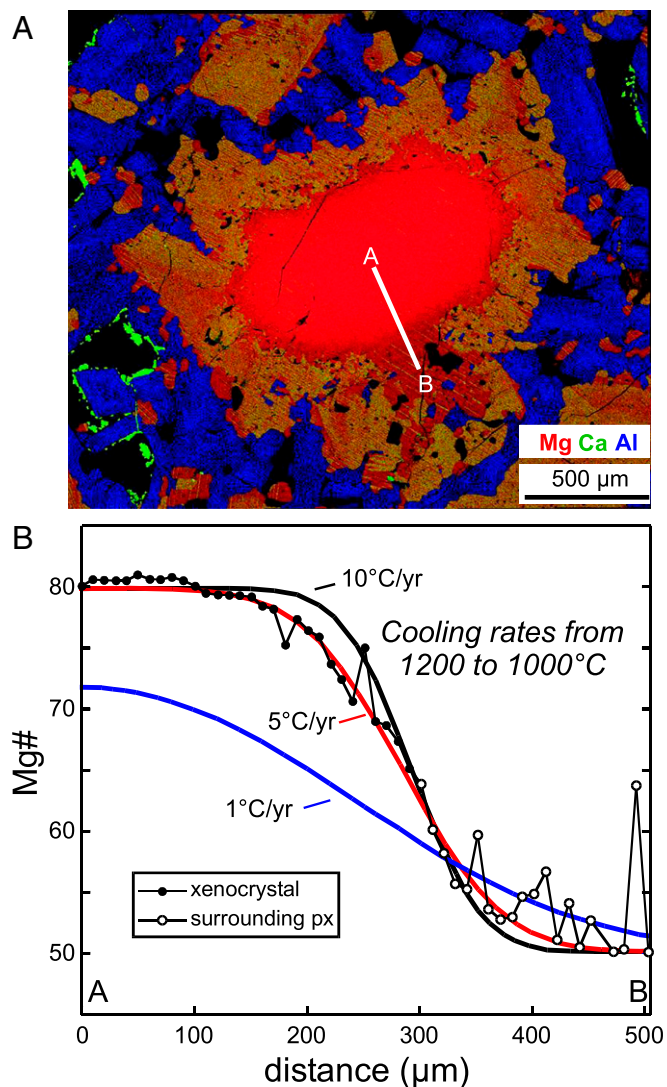
**Fig. 1.** (A) Backscattered electron image of a polished section of EC 002 (pyroxenes are gray to light gray, and feldspars are dark gray). (B) A false-colored X-ray map of the same area. A Mg-rich orthopyroxene xenocryst (red) is seen in the left side of the image.

**Phase Compositions.** Bulk groundmass pyroxene compositions are generally augitic ( $\text{Wo}_{27.7-33.0}\text{En}_{36.9-38.5}\text{Fs}_{30.1-31.0}$ ) (*SI Appendix, Figs. S3–S7 and Tables S1–S4*). Some crystals exhibit remnant zoning from core to rim ( $\text{Wo}_{22.9}\text{En}_{39.4}\text{Fs}_{37.7}$  to  $\text{Wo}_{35.2}\text{En}_{36.3}\text{Fs}_{28.5}$ ). The crystals are partly equilibrated and are a mixture of augite ( $\text{Wo}_{38.7-40.9}\text{En}_{39.5-41.5}\text{Fs}_{18.8-20.7}$ ) and fine exsolutions of low-Ca pyroxene ( $\text{Wo}_{2.4-4.3}\text{En}_{47.1-49.4}\text{Fs}_{47.3-49.3}$ ). Feldspars are albitic ( $\text{Or}_{2.0-7.1}\text{Ab}_{75.8-87.4}\text{An}_{6.7-21.6}$ ) and contain lamellae of K-rich feldspar ( $\text{Or}_{84.0-84.4}\text{Ab}_{11.3-11.6}\text{An}_{4.3-4.4}$ ). Groundmass spinel compositions are  $\text{Usp}_{18.2-56.5}\text{Sp}_{2.6-6.2}\text{Cm}_{94.2-96.5}$  and  $\text{Mg}\# = [100 \times \text{Mg}/(\text{Mg} + \text{Fe}), \text{atomic}] = 3.51$  to  $5.83$  with no Zn ( $<0.02$  wt%). Silica phases contain detectable amounts of Al, Fe, Ca, and alkalis ( $\text{Al}_2\text{O}_3 = 1.8$  to  $2.7$  wt%;  $\text{K}_2\text{O} < 0.5$  wt%;  $\text{Na}_2\text{O} = 0.7$  to  $1.4$  wt%;  $\text{CaO} = 0.03$  to  $0.15$  wt%; and  $\text{FeO} < 0.3$  wt%).

We analyzed two xenocrysts: fragments of a 1-cm-long olivine megacryst and a small pyroxene. The olivine is forsteritic ( $\text{Fo}_{88}$ ,  $\text{FeO}/\text{MnO} = 24.2$  wt%/wt%), Ca rich ( $\text{CaO} = 1.3$  wt%), and poor in Ni ( $1.3$   $\mu\text{g/g}$ ) and Co ( $10.2$   $\mu\text{g/g}$ ). Its light rare earth element (REE) and alkali abundances are much higher than expected ( $\text{La} = 11.7$  ng/g,  $\text{K} = 102$   $\mu\text{g/g}$ ,  $\text{Rb} = 4.9$   $\mu\text{g/g}$ , and  $\text{Cs} = 0.28$   $\mu\text{g/g}$ ) and could reflect the presence of melt inclusions. The small pyroxene is a low-Ca pyroxene (Fig. 2). It displays a chemically homogeneous core ( $\text{Wo}_{4.7}\text{En}_{76.8}$ ) with zoned rims, mantled by groundmass pyroxenes.

The cores have relatively high  $\text{Cr}_2\text{O}_3$  ( $\sim 0.5$  wt%) and  $\text{Al}_2\text{O}_3$  ( $\sim 0.3$  wt%) and low  $\text{TiO}_2$  contents ( $\sim 0.1$  wt%) compared to low-Ca pyroxene in the groundmass. The  $\text{FeO}/\text{MnO}$  ratios of the xenocryst core ( $21.1$  wt%/wt%) are identical to those of the low-Ca pyroxene in the rims and in groundmass ( $\sim 20$  to  $21$  wt%/wt%). Pyroxenes surrounding the crystal are more magnesian than those in the groundmass. Spinel grains found inside this xenocryst are more Cr and Al rich than those found in the groundmass ( $\text{Usp}_{0.9}$  to  $_{2.9}\text{Sp}_{12.4}$  to  $_{17.3}\text{Cm}_{84.3}$  to  $_{88.1}$  and  $\text{Mg}\# = 11.9$  to  $15.7$ ).

These olivine or pyroxene xenocrysts could be debris from the nascent mantle of the protoplanet (melting residues) or crystals formed from previous magmas. A definitive answer cannot be given with the study of only two crystals. Note that the largest pyroxene xenocrysts reach 9 cm (14) and are certainly not “mantle like.” Similar  $\text{FeO}/\text{MnO}$  ratio and  $\Delta^{17}\text{O}$  (14) suggest they are more likely derived from the crystallization of associated magma(s) more magnesian than EC 002.

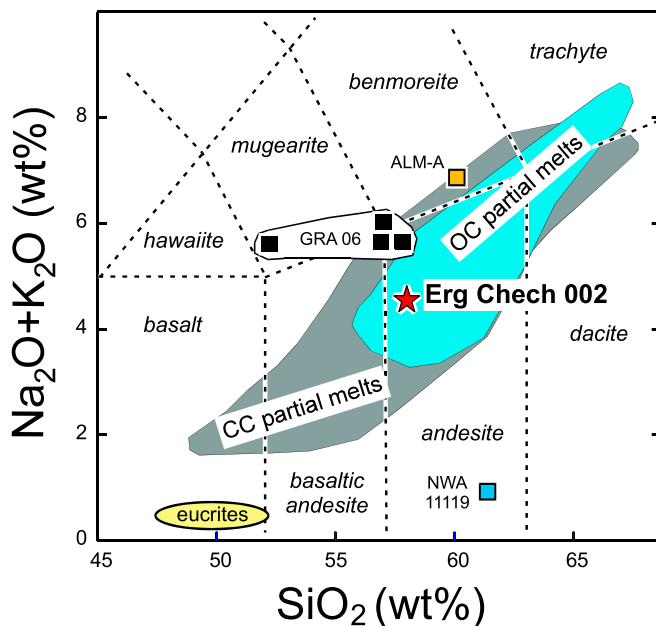


**Fig. 2.** (A) A false-colored X-ray map of a small orthopyroxene xenocryst (in red) mantled by groundmass pyroxenes. (B)  $\text{Mg}\#$  profile across the boundary between xenocryst and groundmass pyroxene. We calculated cooling rates assuming a compositionally homogeneous spherical xenocryst in a homogeneous medium (16). We used the interdiffusion coefficients of Fe-Mg in orthopyroxene ( $\text{Mg}\# = 80$ ) along the c and b directions (17). We chose the initial temperature of  $1,200$   $^{\circ}\text{C}$ , which is within the range of estimated crystallization temperatures and assumed constant cooling rates.

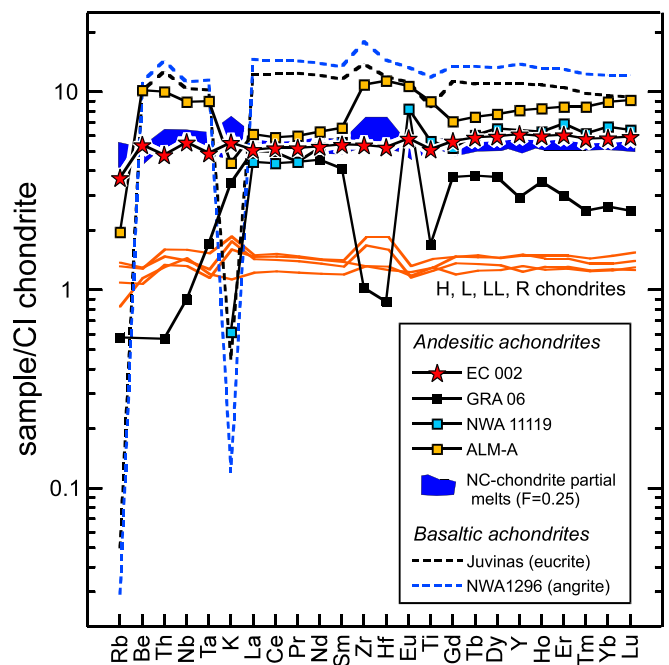
**Geochemistry.** We crushed a 1.1-g sample devoid of apparent xenocrysts, representative of the groundmass, and therefore very close to the melt from which EC 002 crystallized. With 58 wt% SiO<sub>2</sub> and 4.54 wt% of Na<sub>2</sub>O+K<sub>2</sub>O (Fig. 3 and *SI Appendix, Table S4*), it is andesitic according to the International Union of Geological Sciences criteria (18). The rock is quite rich in MgO and FeO. Its CIPW norm indicates proportions of pyroxenes and plagioclase in accordance with modal estimates (53 wt% pyroxene and 40.5 wt% plagioclase [i.e., about 47 vol% pyroxene and 47 vol% plagioclase taking into account the mineral densities]). The calculation also shows some normative quartz (2.5 wt%), indicating a rock slightly oversaturated in SiO<sub>2</sub>. EC 002 is very poor in P (216 μg/g), Ni (18.5 μg/g), Co (5.85 μg/g), Cu (1.4 μg/g), Pb (90 ng/g), W (22 ng/g), Ga (2.6 μg/g), and Zn (0.44 μg/g). EC 002 displays some excesses in Ba, Sr, and U, which are usual for Saharan finds (e.g., ref. 19) and will not be further discussed. Other incompatible trace element abundances (e.g., REE, Th, etc.) are low and of the order of 5 or 6 times the chondritic reference. Indeed, the CI-normalized trace element pattern of EC 002 is rather flat with no noticeable anomaly, even for high field strength elements and alkalis (Fig. 4). This rock is only slightly light REE depleted (La<sub>n</sub>/Sm<sub>n</sub> = 0.94) with a small positive Eu anomaly (Eu/Eu\* = 1.05). Moreover, it displays the same Tm negative anomaly (Tm/Tm\* = 0.973) as noncarbonaceous (NC) chondrites (Tm/Tm\* < 1 and typically 0.97 to 0.98) and achondrites and inner solar system planetary bodies [Vesta, Mars, Moon, and Earth, with average Tm/Tm\* typically ~0.975 (21)].

The chemical composition of EC 002 is very different from that of other andesitic achondrites. It is, for example, much richer in alkalis than NWA 11119, without reaching the concentrations of ALM-A or GRA 06 (Fig. 3). The smoothness of its trace element pattern (Fig. 4), parallel to that of NC chondrites even for alkalis, also distinguishes it from the other known achondritic lavas.

**<sup>26</sup>Al-<sup>26</sup>Mg Systematics.** A total of 19 feldspar grains and 11 pyroxenes were analyzed (*SI Appendix, Tables S5 and S6*). The pyroxenes are characterized by very low <sup>27</sup>Al/<sup>24</sup>Mg ratios ranging from 0.018 to 0.06 and homogeneous (within their typical ±0.15%



**Fig. 3.** A plot of total alkalis versus silica content showing the compositions EC 002, the other andesitic achondrites, eucrites, and the experimental melts obtained on chondritic systems by Collinet and Grove (12).

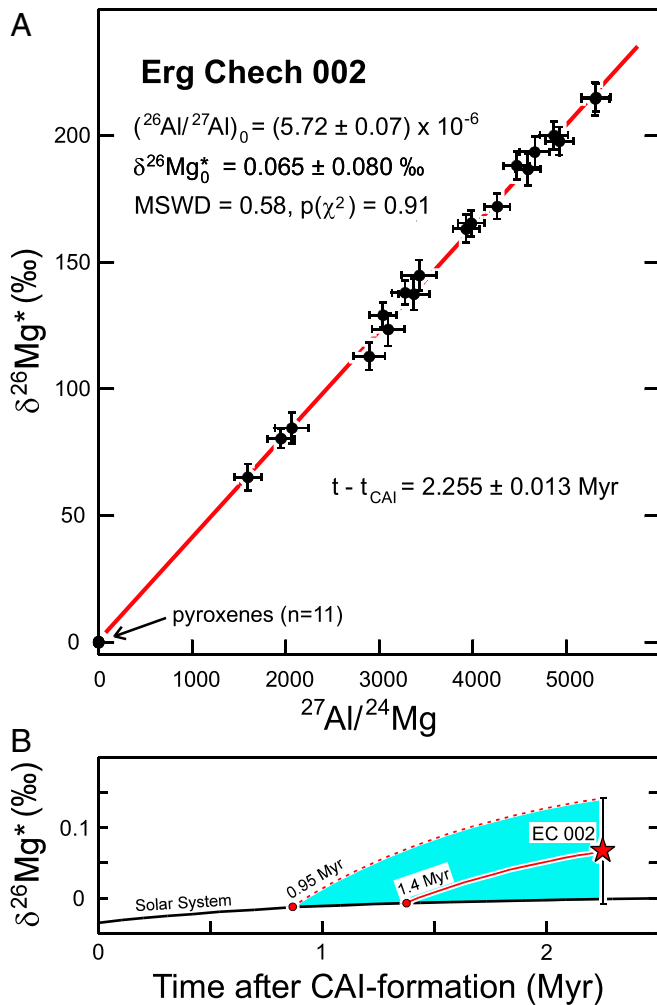


**Fig. 4.** Comparison of CI-normalized trace element abundances in EC 002, other andesitic achondrites, representative basaltic achondrites, and NC chondrites (6–9). The field of partial melts (F = 25%) obtained from regular NC chondrites is shown for comparison. CI normalization values are from ref. 20.

2 SE errors)  $\delta^{26}\text{Mg}^*$  from  $-0.10$  to  $+0.25\text{‰}$  (average  $+0.067 \pm 0.076\text{‰}$  [2SE]). Plagioclases display a wide range of radiogenic Mg enrichments, with  $^{27}\text{Al}/^{24}\text{Mg}$  from 1,597 to 5,316 and  $\delta^{26}\text{Mg}^*$  from 65 to 215‰. Pyroxenes and feldspar define a  $^{26}\text{Al}$  isochron (Fig. 5) with an initial  $^{26}\text{Al}/^{27}\text{Al}$  ratio of  $(5.72 \pm 0.07) \times 10^{-6}$ , the highest value ever reported for an achondrite. This ratio translates into a closure age of the  $^{26}\text{Al}$ - $^{26}\text{Mg}$  system of 4,565.0 Myr (i.e.,  $2,255 \pm 0.013$  Myr after Ca-Al-rich inclusions [CAIs] assuming a canonical distribution of  $^{26}\text{Al}$  (22), or of 4,566.1 Myr (i.e., 1 Myr after CAIs) using the D’Orbigny angrite anchor (23, 24). The calculated initial  $\delta^{26}\text{Mg}^*$  ( $\delta^{26}\text{Mg}^*_0 = +0.065 \pm 0.08\text{‰}$  [2SE]) is consistent with the average composition of the pyroxenes and significantly higher than the canonic initial of the solar system of  $-0.040 \pm 0.029\text{‰}$  as defined from CAIs (22).

**Noble Gas Isotope Composition.** Abundances and isotope ratios of He, Ne, and Ar were analyzed by step heating (*SI Appendix, Table S7*). Isotope ratios of He and Ne are strongly enriched in cosmogenic isotopes, with concordant  $^3\text{He}$  and  $^{21}\text{Ne}$  exposure ages of  $26.0 \pm 1.6$  Myr and  $25.6 \pm 1.0$  Myr, respectively. The bulk  $^{40}\text{Ar}/^{36}\text{Ar}$  ratio ( $5,522 \pm 42$ ) is strongly enriched in radiogenic  $^{40}\text{Ar}$  from the decay of  $^{40}\text{K}$ . The  $^{40}\text{Ar}$  abundance corresponds to a K-Ar age of  $4534^{+117}_{-125}$  Myr, within uncertainty of the closure age for the  $^{26}\text{Al}$ - $^{26}\text{Mg}$  system. This indicates that Ar has been effectively retained within EC 002 since shortly after formation, precluding significant thermal events and implying a relatively early breakup of the parent body. Furthermore, the absence of significant trapped “planetary” noble gas components suggests that the parent melt of EC 002 was efficiently degassed prior to crystallization. After correction for cosmogenic  $^{36}\text{Ar}$ , trapped  $^{36}\text{Ar}$  abundance is only  $4.3 \times 10^{-13} \text{ mol} \cdot \text{g}^{-1}$ , which is more than an order of magnitude lower than typical achondrite abundances (25).

**Spectroscopy.** Reflectance spectra of EC 002 reveal the presence of two strong absorptions related to Ca-rich pyroxene. This spectral signature does not correspond to any known asteroid spectral type.



**Fig. 5.** (A) The Al-Mg internal isochron defined by feldspars and pyroxenes of EC 002. The error bars correspond to  $2 \times SEs$  on the ratios. (B) Mg isotopic composition of EC 002 compared to the theoretical evolution of the Mg isotopic composition of the solar system [calculated for a chondritic  $^{27}Al/^{24}Mg$  ratio of 0.101, a  $(^{26}Al/^{27}Al)_0$  of  $5.23 \times 10^{-5}$ , and a  $\delta^{26}Mg^*$  of  $-0.034\text{‰}$  (22)].

Comparison between colors of EC 002 and those from about 10,000 objects from the Sloan Digital Sky Survey database also reveal the rarity if not absence of similar objects within the asteroid population (*SI Appendix, section B*).

## Discussion

As demonstrated by its chemistry and Al-Mg age, EC 002 is a unique fragment of the crust of an ancient, differentiated body, contemporaneous with the formation of the cores of the parent bodies of iron meteorites (26). The oldest igneous rock previously described was NWA 11119 (9), with an age 1.24 Myr younger than EC 002 when calculated with the same canonical compositions. For major elements, EC 002 is chemically distinct from all the other basaltic, andesitic, or trachyandesitic achondrites (Fig. 3). Its alkali to refractory incompatible element ratios is unfractionated, making EC 002 very different from the crust of telluric planets (Fig. 4) in the sense that it is not depleted in volatile elements. For example, its K/Th ratio ( $=22,500$ ) is similar to that of CI chondrites [ $=19,400$  (20)] and much higher than that of the crusts of Mercury, Earth, or Mars ( $\sim 5,500$  or below), as well as the Moon, Vesta, and the parent body of the angrites, which are strongly alkali depleted (27, 28). However, its very low Zn and

Ga abundances point to a possible depletion for some of the other volatile elements, inherited from its accretionary materials. These low concentrations could also be the fingerprints of the early segregation of some metal and sulfides (initiation of the core formation), thus explaining the low abundances of Ni, Co, and W. EC 002 displays a Tm/Tm\* ratio identical to those of differentiated bodies from the inner solar system and clearly points to a NC affinity (21). Thus, despite Ti, Cr, or Mo isotopic data not yet being available, the parent body of EC 002 can most likely be affiliated to the NC family of objects formed early in the accretion disk (e.g., refs. 29 and 30). Oxygen isotopic composition of EC 002 is consistent with this interpretation. It is similar to that of some ungrouped basaltic (NC) achondrites such as Bunburra Rockhole, Asuka 881394, and Emmaville, but which are mineralogically and chemically too different to originate from the same parent body as EC 002 (14).

The genesis of EC 002 allows us to understand some aspects of primordial crust formation on bodies of chondritic compositions. Though an andesite can derive from the fractional crystallization of a more primitive melt, this cannot be the case for EC 002. First, EC 002 has low abundance of highly incompatible elements (e.g., REE or Th). If it were an evolved lava, its parental melt would have even lower concentrations, which is difficult to envisage even for a primary basalt. The concentration of MgO (7.06 wt%) and the Mg#-number ( $= 52.9$ ) of EC 002 are quite high for an andesite and do not support the idea that significant amounts of ultramafic cumulates were extracted. Furthermore, fractional crystallization would have inevitably left its fingerprint in the distribution of trace elements, such as a marked negative Eu anomaly generated by plagioclase crystallization at variance with the composition of EC 002 (Fig. 4). Therefore, it must be considered that EC 002 could be a primitive or even a primary melt. Partial melting of chondrites at low pressure is perfectly capable of generating andesitic magmas rich in  $SiO_2$  and alkalis (10–12). However, the major element composition of EC 002 lies at the high-melting end of the experimental melting trends defined for ordinary H or LL chondrites (Fig. 3 and *SI Appendix, Fig. S8*). These trends indicate that melts with the same composition as EC 002 are obtained after plagioclase exhaustion for high degrees of melting (F) of around 25% (*SI Appendix, Fig. S9*). Such high melting rates would also explain the unfractionated trace element pattern of EC 002 since all phases with high crystal-melt partition coefficients (such as phosphates or plagioclase) would be exhausted in the source after  $\sim 17\%$  partial melting (12, 31). Indeed, the trace element pattern of EC 002 fits with an enrichment by a factor of  $\sim 1/F$  of a typical NC chondrite composition (Fig. 4), in perfect agreement with the experimental data.

The thermal history of EC 002 appears quite straightforward. Experimental data allow a precise evaluation of the temperature at which the magma formed (*SI Appendix, Fig. S10*). The MgO of EC 002 gives an estimated melting temperature of  $1,224 \pm 20$  °C. This high temperature is confirmed by the crystallization temperatures estimated from the bulk compositions of Ca-rich pyroxene (32) in the range of 1,149 to 1,229 °C (average = 1,186 °C [SD = 25 °C], slightly lower due to postcrystallization equilibration during cooling). The final equilibration temperature, about 957 °C, is estimated from the highest Ca concentration in pyroxene (32). The cooling of the rock was fast enough to preserve remnant zoning in groundmass pyroxene and the compositions of the core of the small xenocrysts. As a demonstration, we tentatively modeled the zoning profile of Mg# across the small xenocryst exposed in our section (Fig. 2 and *SI Appendix*). The cooling rate is estimated to be about 5 °C/y between 1,200 and 1,000 °C, a value consistent with a thick lava flow or a shallow intrusion. Important additional constraints are brought by silica polymorphs (*SI Appendix*). Only cristobalite and tridymite (PO) were detected in EC 002, and quartz is totally lacking. As experimentally shown for eucrites (33, 34), cristobalite crystallized at high temperatures.

Subsequently, it partially transformed to tridymite above  $\sim 900$  °C. Since cristobalite easily transforms to quartz (34), the lack of quartz indicates a very fast cooling rate below 900 °C ( $>0.1$  to  $1$  °C/d), consistent with the absence of monoclinic tridymite and the possible occurrence of anorthoclase. The most likely explanation for this change in cooling rate is an impact that would have excavated, or more likely ejected, the rock from its parent body, in agreement with the evidence for shock metamorphism (M-S2). The similarity between the K-Ar retention age and the formation age as calculated from Al-Mg indicates that EC 002 was not significantly heated following formation, further supporting the idea that any ejection event occurred rapidly after formation. The cooling history of this meteorite therefore appears to be short, since it would have cooled for only a few decades before it was probably ejected.

The Mg isotopic composition of EC 002 gives further clues on the timing of differentiation of its parent body. Because of the fast cooling inferred for EC 002, the old  $^{26}\text{Al}$  age of  $2.255 \pm 0.013$  Myr after CAIs can be considered to date the crystallization of the parent melt. This age is tightly constrained from the following: 1) the quality of the  $^{26}\text{Al}$  isochron (Fig. 5A) that also implies a lack of significant  $^{26}\text{Mg}$  redistribution by metamorphism after this fast cooling and 2) the fact that EC 002 is related to inner solar system NC bodies which are considered to have formed with a canonic level of  $^{26}\text{Al}$  even in the hypothesis of a heterogeneous distribution of  $^{26}\text{Al}$  in the accretion disk (35, 36). The fact that the  $^{26}\text{Al}$  isochron intercept gives a  $\delta^{26}\text{Mg}^*_0$  significantly higher than the canonic initial of the solar system (Fig. 5B) is likely an indication for a protracted history of the parental melt of EC 002 prior to crystallization. In fact, such excesses of  $^{26}\text{Mg}$  have not been observed for bulk chondrites (e.g., refs 37 and 38) but are known in two ancient achondrites [NWA 7325:  $\delta^{26}\text{Mg}^*_0 = 0.093 \pm 0.004\%$  (39); Asuka 881394:  $\delta^{26}\text{Mg}^*_0 = 0.070 \pm 0.052\%$  (40)]. They could result from metamorphic perturbations or redistributions (40) or correspond to the protolith isotopic composition in the case of remelting of a crustal reservoir (39, 41). Here, these explanations are not satisfactory due to the short cooling duration of EC 002 and the evidence that the andesitic magma formed directly from a chondritic source. In addition, because the olivine and pyroxene xenocrysts present in EC 002 have a very low Al/Mg ratio (and thus cannot develop significant radiogenic  $^{26}\text{Mg}$  excesses), their eventual assimilation by the EC 002 parental melt is also unable to explain the positive  $\delta^{26}\text{Mg}^*_0$ . Alternatively, since EC 002 has a superchondritic Al/Mg ratio, a high  $\delta^{26}\text{Mg}^*_0$  value can simply be acquired if the time elapsed between partial melting and crystallization is long enough for  $^{26}\text{Mg}$  radiogenic excesses to develop in the melt. In order to investigate this possibility further, the Mg isotopic evolutionary curves of EC 002 and the solar system (equivalent to the evolution of regular chondrites) were compared (Fig. 5B). The two curves intersect between 0.95 Ma and 2.25 Ma (the age of crystallization) after the formation of CAIs. While this range is wide due to the error the  $\delta^{26}\text{Mg}^*_0$  value of EC 002, it shows that the transfer of magmas between the melting zones and the surface could have been slow, a duration exceeding a few  $10^5$  y being probable. This result confirms and strengthens the inferences made by Collinet and Grove (12), who calculated that the velocity extraction of silica- and alkali-rich magmas in a protoplanet could have been at least three orders of magnitude lower than that of basalts (2.3 to 230 m/My versus 1.1 to 1,100 km/My, respectively). Thus, these magmas would have moved slowly, probably only over narrow distances during one half-life of  $^{26}\text{Al}$ , rendering the thickening of a primordial crust by the stacking of flows difficult.

The parent body of EC 002 was certainly not unusual, and despite the scarcity of andesitic achondrites identified to date, it is reasonable to assume that many similar chondritic bodies accreted at the same time and were capped by the same type of primordial crust as proposed by Collinet and Grove (12). However, this

andesitic crust might have been only temporary. Thermal models of asteroids which undergo partial melting due to the heat generated by  $^{26}\text{Al}$  decay (e.g., refs. 26 and 42) show that if accretion starts before 1 Myr after CAIs, the average temperature of the body reaches at least  $1,500$  °C at 2.2 Myr and can continue to rise after that. This is true whether accretion takes place instantaneously by gravitational instability or at a slower pace by a combination of gravitational instability and pebble accretion. Such temperatures are incompatible with the thermal history reconstructed for EC 002. One solution to this problem would be to consider the following: 1) that the parent body of EC 002 accreted later than 1 Myr (i.e., around 1.5 Myr) and 2) that EC 002 formed at the surface or close to the surface of the parent body (as implied by the texture and cooling history of EC 002). Such a body could escape global melting and have melts generated at  $\sim 1,220$  °C close to surface migrating over a few 100 kyr to the surface. This would allow the preservation of a primordial crust at the surface of the parent body of EC 002. Such andesitic crusts were perhaps quite common 4,565 Myr ago. Possible parent bodies can be sought among available observations of asteroid surfaces. The spectral data obtained on EC 002 were compared with those obtained on asteroids, taking into account space weathering, the presence of olivine xenocrysts, and residual mantle debris (*SI Appendix*). EC 002 is clearly distinguishable from all asteroid groups (43, 44), and no object with spectral characteristics similar to EC 002 has been identified to date. Remains of primordial andesitic crust are therefore not only rare in the meteorite record, but they are also rare today in the asteroid belt. This suggests that the earliest differentiated protoplanets that populated the solar system, as well as most of their debris, were certainly destroyed or subsequently accreted to the growing rocky planets, making the discovery of meteorites originating from primordial crusts an exceptional occurrence.

## Materials and Methods

**Petrography and Phase Compositions.** We studied two polished sections ( $\sim 1.5$  cm<sup>2</sup> each), one thick, one thin, and made from the same slab. We examined these sections using an optical microscope equipped with both transmitted and reflected light, an electron microprobe analyzer (EPMA; JEOL JXA-8200), and a field emission scanning electron microscope (JEOL JSM-7100F) equipped with an energy dispersive spectrometer (Oxford AZtec Energy) at the National Institute of Polar Research (NIPR). Pyroxene and spinel were analyzed using a current of 30 nA and a focused beam and plagioclase using 10 nA and a defocused beam ( $\sim 3$   $\mu\text{m}$  in diameter) at 15 keV. For Mn analysis, we used a LIFH spectrometer. Count rates using the LIFH were increased by a factor of 3 to 4 compared to those of LIF. Bulk compositions of pyroxene are obtained from averaging 10 to 20 points across the grains using a beam diameter of  $\sim 30$   $\mu\text{m}$  at 15 keV. Data were reduced using a ZAF correction program. Mineral phases were identified with an inVia Raman spectrometer at NIPR.

**Geochemistry.** Another slice weighing 1.09 g was chosen after a thorough binocular examination of both sides because it contained no apparent pyroxene or olivine xenocrysts. It was crushed using a boron carbide mortar and pestle into a homogeneous fine-grained powder in clean room conditions at Institut Universitaire Européen de la Mer, Plouzané. Clear fragments (45 mg) of a large ( $>1$  cm) olivine crystal were leached in 6 N HCl (120 °C, 30 min), rinsed three times, and dried. Major and trace elements were determined respectively by inductively coupled plasma-atomic emission spectroscopy using a Horiba Jobin Yvon Ultima 2 spectrometer, and by inductively coupled plasma-sector field mass spectrometry using a Thermo Scientific ELEMENT XR spectrometer. We used the same procedures as Cotten et al. (45) for the major elements. The accuracy is better than 5% for Na and P and much better than 3% for the other elements. For trace elements, we used the same procedure as Barrat et al. (20, 21). Based on standards and many sample duplicates, the precisions for abundances are in most cases much better than 5% [two relative SDs ( $2 \times \text{RSD}$ )]. The precisions for Eu/Eu\* and Tm/Tm\* ratios [ $X^*$  is the expected X concentration for a smooth CI-normalized REE pattern, such that  $\text{Eu}_n = (\text{Sm}_n \times \text{Gd}_n)^{1/2}$  and  $\text{Tm}_n^* = (\text{Er}_n \times \text{Yb}_n)^{1/2}$ ] are respectively better than 3% and 1.5% ( $2 \times \text{RSD}$ ).

**$^{26}\text{Al}$ - $^{26}\text{Mg}$  Dating.** A thick slice (1 cm<sup>2</sup>) was polished and mounted in a four-window ion probe holder. The Mg isotopic compositions and Al/Mg concentration ratios were measured with the CRPG-CNRS CAMECA ims 1280-HR2 ion microprobe.

**Ion probe settings for feldspar.** Because of the low Mg contents (<0.03 wt% of MgO) of the feldspars, the measurements were made in monocollection mode using the central electron multiplier (EM) for  $^{24}\text{Mg}^+$ ,  $^{25}\text{Mg}^+$ , and  $^{26}\text{Mg}^+$  and the central Faraday cup (FC2) for  $^{27}\text{Al}^+$ . The samples were sputtered with a  $\sim 5$  nA  $\text{O}^-$  primary beam rastered at  $15 \times 15 \mu\text{m}$ . The transfer optic magnification was set at 100  $\mu\text{m}$  to ensure an efficient instrumental transmission and to fully fill the field aperture set at 2,000  $\mu\text{m}$ . The mass resolving power (MRP) was set at  $M/\Delta M \sim 5,000$  in order to completely remove the  $^{24}\text{MgH}^+$  interference on  $^{25}\text{Mg}^+$ . One measurement consisted of a 120 s preanalysis sputtering to clean the sample surface and attain stable count rates on detectors followed by automatic secondary beam and energy centering, and 40 cycles with counting times during each cycle of 4, 5, 10, 10, 3, and 3 s at masses 23.8 (background for EM),  $^{24}\text{Mg}$ ,  $^{25}\text{Mg}$ ,  $^{26}\text{Mg}$ , 26.8 (background FC2,) and  $^{27}\text{Al}$ , respectively. The Mg isotopic compositions are given in delta notation according to  $\delta^{25}\text{Mg}_x = \ln[(^{25}\text{Mg}/^{24}\text{Mg})_x / (^{25}\text{Mg}/^{24}\text{Mg})_{\text{standard}}] \times 1,000$  (similarly for  $^{26}\text{Mg}$ ) with  $(^{25}\text{Mg}/^{24}\text{Mg})_{\text{standard}} = 0.12663$  and  $(^{26}\text{Mg}/^{24}\text{Mg})_{\text{standard}} = 0.13932$  (46). The  $^{26}\text{Mg}$  excesses due to  $^{26}\text{Al}$  decay are noted  $\delta^{26}\text{Mg}^*$  with  $\delta^{26}\text{Mg}^* = \delta^{26}\text{Mg} - \delta^{25}\text{Mg}/\beta$  [with  $\beta = 0.521$  for an equilibrium mass fractionation of Mg isotopes as it is expected for the current object (47)]. The choice of the mass fractionation law is not critical to the current data set since the degree of intrinsic mass fractionation is small and most likely due to imperfectly corrected matrix effects in feldspars (SI Appendix, Table S5), and because the Al/Mg ratios of feldspar that control the isochron are high enough for the uncertainty introduced by the choice of fractionation law to be insignificant. Miyake-Jima plagioclase ( $^{27}\text{Al}/^{24}\text{Mg} = 396.3$ ) was used to calibrate the instrumental isotopic fractionation and the relative Al/Mg ion yield. Two sigma SEs on the mean of  $\pm 0.40\%$ ,  $\pm 0.31\%$ , and  $\pm 0.45\%$  for  $\delta^{25}\text{Mg}$ ,  $\delta^{26}\text{Mg}$ , and  $\delta^{26}\text{Mg}^*$ , respectively, were obtained for the standard. The relative Al/Mg ion yield was determined to be  $0.833 \pm 0.028$ . Typical counts rates for EC 002 feldspars were 15,000 cps, 2,000 cps, 2,500 cps, and  $4.2 \times 10^7$  cps  $^{24}\text{Mg}$ ,  $^{25}\text{Mg}$ ,  $^{26}\text{Mg}$ , and  $^{27}\text{Al}$ , respectively. The errors reported for the measurements of the samples are two sigma errors calculated by summing in a quadratic way the errors due to counting statistic in each point and the errors due to calibration of instrumental isotopic fractionation and Al/Mg ion yield (SI Appendix, Table S5).

**Ion probe settings for pyroxene.** The measurements were made in multi-collection mode using four off-axis FCs (L1, C, H1, and H'2 for  $^{24}\text{Mg}^+$ ,  $^{25}\text{Mg}^+$ ,  $^{26}\text{Mg}^+$ , and  $^{27}\text{Al}^+$ , respectively). The samples were sputtered with a  $\sim 6$  nA  $\text{O}^-$  primary beam rastered at  $15 \times 15 \mu\text{m}$ . The transfer optic magnification was set at 80  $\mu\text{m}$  to insure an efficient instrumental transmission and to fully fill the field aperture set at 2,500  $\mu\text{m}$ . The MRP was set at  $M/\Delta M \sim 5,000$  (slit 2) in order to completely remove the  $^{24}\text{MgH}^+$  interference on  $^{25}\text{Mg}^+$ . One measurement consisted of a 90-s preanalysis sputtering to clean the sample surface and attain stable count rates on detectors, during which offsets of FCs were measured, followed by automatic secondary beam and energy centering, and 40 cycles of 5-s integration time for data acquisition. Gold enstatite ( $^{27}\text{Al}/^{24}\text{Mg} = 0.024$ ) was used to calibrate the instrumental isotopic fractionation and the relative Al/Mg ion yield. Two sigma SEs on the mean of  $\pm 0.04\%$ ,  $\pm 0.05\%$ , and  $\pm 0.04\%$  for  $\delta^{25}\text{Mg}$ ,  $\delta^{26}\text{Mg}$ , and  $\delta^{26}\text{Mg}^*$ , respectively, were obtained for the standard. The relative Al/Mg ion yield was determined to be  $0.788 \pm 0.009$ . The errors reported for the measurements of the samples are two sigma errors calculated by summing in a quadratic way the errors due to counting statistic in each point and the errors due to calibration of instrumental isotopic fractionation and Al/Mg ion yield (SI Appendix, Table S6).

The  $(^{26}\text{Al}/^{27}\text{Al})_0$  and  $\delta^{26}\text{Mg}^*_0$  were calculated from the isochron regression fitted using Isoplot software. The  $^{26}\text{Al}$ - $^{26}\text{Mg}$  relative age was calculated using the canonical  $^{26}\text{Al}/^{27}\text{Al} = 5.23 \times 10^{-5}$  (22) and a  $^{26}\text{Al}$  half-life of 705 kyr (48).

**Noble Gas Isotope Analysis.** An 8.2-mg piece of the sample was loaded into a filament furnace consisting of three alumina-coated tungsten evaporation baskets. The sample was placed directly into one of the baskets while two empty baskets served to calculate the blank contribution. The full analytical procedure has been previously documented in Broadley et al. (49). In brief, the furnace was pumped while baking at 150  $^\circ\text{C}$  for 48 h to remove adsorbed atmospheric gases from the walls of the furnace and the sample. The furnace was then pumped for a further 48 h to ensure low blank levels. Gases were extracted from the sample over five temperature steps at 600  $^\circ\text{C}$ , 800  $^\circ\text{C}$ , 1,000  $^\circ\text{C}$ , 1,200  $^\circ\text{C}$ , and 1,400  $^\circ\text{C}$ . The majority of the He (90%) was released at the 800  $^\circ\text{C}$  extraction step. For both Ne (84%) and Ar (91%), the majority was released at 1,200  $^\circ\text{C}$ . Negligible amounts of gas were released at 1,400  $^\circ\text{C}$ , and inspection of the sample after analysis confirmed it had been completely vaporized.

The extracted He, Ne, and Ar were purified, cryo-separated, and analyzed using a Helix-MC mass spectrometer (49). Blanks were analyzed following exactly the same protocol as samples, and for the major extraction step, the  $^4\text{He}$ ,  $^{21}\text{Ne}$ , and  $^{40}\text{Ar}$  blanks represent 0.01%, 0.01%, and 0.14% of the sample, respectively. Blank contributions for  $^{36}\text{Ar}$  were significantly higher given the very low concentration of  $^{36}\text{Ar}$  in the sample, with blank contribution reaching a maximum of 50% for the 1,000  $^\circ\text{C}$  temperature step. Noble gas abundances and isotope ratios are reported in SI Appendix, Table S7.

Exposure ages were calculated for  $^3\text{He}$  and  $^{21}\text{Ne}$  following Leya et al., (50) using target element abundances from the bulk chemical composition shown in SI Appendix, Table S4 and a shielding depth correction factor of 20 cm. The K-Ar age was calculated following Kelley (51), assuming that 100% of the measured  $^{40}\text{Ar}$  is radiogenic. A correction for atmospheric  $^{40}\text{Ar}$  results in a change of <1% and is insignificant compared to the uncertainty already associated with the measurement. The trapped  $^{36}\text{Ar}$  component was calculated using simple 2-endmember mixing, using a theoretical cosmogenic  $^{38}\text{Ar}/^{36}\text{Ar}$  of 1.438 and an assumed trapped  $^{38}\text{Ar}/^{36}\text{Ar}$  of 0.188 [atmospheric but also similar to other trapped planetary phases (52)]. This gives the fraction of trapped  $^{36}\text{Ar}$  as 21% and a total trapped abundance of  $4.3 \times 10^{-13}$  mol/g.

**Reflectance Spectra.** Reflectance spectra of Erg Chech 002 were measured at Institut de Planétologie et d'Astrophysique de Grenoble using the Shadows instrument (53). We used the standard mode of the instrument (around 7-mm diameter illumination spot), and spectra were measured under nadir illumination and using an observation angle of 30 $^\circ$ . One spectrum was obtained for a powdered sample, and three spectra were obtained on distinct location of a raw slab of the meteorite.

**Data Availability.** All study data are included in the article and/or SI Appendix.

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