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PLANETARY SCIENCE

Oxygen isotopes of anhydrous primary minerals show kinship between asteroid Ryugu and comet 81P/Wild2

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The extraterrestrial materials returned from asteroid (162173) Ryugu consist predominantly of low-temperature aqueously formed secondary minerals and are chemically and mineralogically similar to CI (Ivuna-type) carbonaceous chondrites. Here, we show that high-temperature anhydrous primary minerals in Ryugu and CI chondrites exhibit a bimodal distribution of oxygen isotopic compositions: ¹⁶O-rich (associated with refractory inclusions) and ¹⁶O-poor (associated with chondrules). Both the ¹⁶O-rich and ¹⁶O-poor minerals probably formed in the inner solar protoplanetary disk and were subsequently transported outward. The abundance ratios of the ¹⁶O-rich to ¹⁶O-poor minerals in Ryugu and CI chondrites are higher than in other carbonaceous chondrite groups but are similar to that of comet 81P/Wild2, suggesting that Ryugu and CI chondrites accreted in the outer Solar System closer to the accretion region of comets.

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INTRODUCTION

Samples returned from asteroid (162173) Ryugu by the JAXA Hayabusa2 spacecraft (1) have mineralogical, petrological, and chemical characteristics similar to those of CI (Ivuna-type) carbonaceous chondrites (2, 3). The Ryugu samples and CI chondrites consist mainly of secondary minerals (phyllosilicates, carbonates, magnetite, and pyrrhotite) aqueously altered from anhydrous primary minerals in their parent bodies at low temperature. High-temperature anhydrous primary minerals, including olivine, low-Ca pyroxene, spinel, hibonite, and perovskite, are rare (2–8). Most primary phases in Ryugu and CI chondrites are small (<~20 μm) and monomineralic (5), providing little information on their origin. The morphology and chemical compositions of olivine grains in Ryugu (2) and CI chondrites (4, 6) are consistent with originating as chondrule phenocrysts. A porous olivine-diopside object in Ryugu (2, 9) could be genetically related to amoeboid olivine aggregates (AOAs), a common type of refractory inclusions observed in chon-

Oxygen isotopic compositions of primary minerals can potentially provide important constraints on their origin. In

carbonaceous chondrites, the O isotopic compositions of minerals in chondrules and refractory inclusions [AOAs and Ca-Al–rich inclusions (CAIs)] show a bimodal distribution of Δ^{17} O, a deviation from the terrestrial mass-dependent fractionation law {= δ^{17} O $-0.52\times\delta^{18}$ O, where δ^{i} O = [(i O/ 16 O)_{sample}/(i O/ 16 O)_{SMOW}-1] \times 1000, i = 17 or 18, and SMOW is standard mean ocean water} (11, 12). Most refractory inclusions have solar-like 16 O-rich compositions with Δ^{17} O of \sim 23 per mil (%o) (12), while chondrules are 16 O-depleted to various degrees with Δ^{17} O values that range from \sim 7 to \sim 0%o (Fig. 1, A to D) (13–17).

The previously reported O isotopic compositions of olivine and low-Ca pyroxene grains separated from CI chondrites (4, 6) show $\Delta^{17}{\rm O}$ values that range from \sim -6 to +3% with a clear mode in the distribution at \sim 0% (Fig. 1E), which is different from the modes of olivine and low-Ca pyroxene chondrule phenocrysts in other carbonaceous chondrites, at \sim -6 and \sim -2%, respectively (Fig. 1, A to D) (13–17). The O isotopic compositions of olivine and low-Ca pyroxene grains embedded in matrices of CI chondrites were measured in situ (7). The data of $^{16}{\rm O}$ -poor grains follow a near-uniform distribution ranging between $\Delta^{17}{\rm O}$ of \sim -7

and -1% without any distinct peak (Fig. 1F). So far, no clear evidence for the genetic relationship between olivine and low-Ca pyroxene grains of CI and chondrule minerals of carbonaceous chondrites has been established. Some $^{16}\text{O-rich}$ ($\Delta^{17}\text{O}$ of \sim -20‰) olivine and low-Ca pyroxene grains were observed in CI chondrites and Ryugu samples (Fig. 1, F and G), possibly related to AOAs (7–9). Here, we report O isotopic compositions of primary minerals (olivine, low-Ca pyroxene, and spinel) measured in situ in polished sections of Ryugu and Ivuna, and we identify the first unambiguous AOA object in CI chondrites. We also discuss the implications of these data for understanding (i) the origin of anhydrous primary minerals in Ryugu and CIs, (ii) the genetic relationship between Ryugu and CI chondrites, and (iii) the accretion region of Ryugu and CI chondrite parent bodies.

RESULTS

Primary minerals, like olivine, low-Ca pyroxene, and spinel, are rare in the Ryugu and Ivuna samples studied, consistent with the previous results (2, 5). They occur primarily in ~100- to 500-µm-sized clasts enriched in S and Fe and depleted in Mg and Si compared to the major lithologies of Ryugu (Fig. 2 and figs. S1 to S3) and Ivuna (figs. S4 and S5). Different chemical compositions of the less-altered clasts from major lithologies may be due to difference in chemical compositions of aqueous fluids (2). The anhydrous primary minerals are embedded in hydrated matrix composed of phyllosilicates (serpentine and saponite) and coarser-grained magnetite, sulfides, and carbonates (Fig. 2 and figs. S6 to S8). The carbonates are almost exclusively calcite; dolomite and breunnerite, commonly observed in the main lithologies of Ryugu and Ivuna, are nearly absent in the Fe-rich clasts. Most olivine grains are

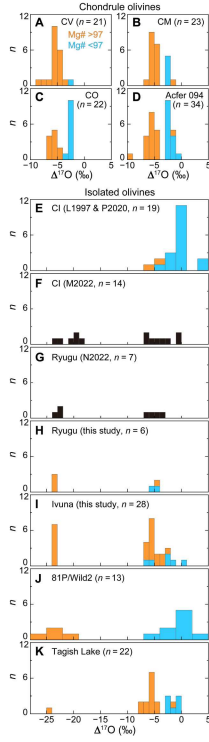
irregularly shaped fragments with sizes up to ~15 μ m in Ryugu and up to ~30 μ m in Ivuna. The chemical compositions of the olivine grains range from Mg# [= Mg/(Mg + Fe) × 100] ~57 to ~99, but the very Mg-rich olivine grains (Mg# > 97) are dominant (fig. S9).

A Ryugu piece from the first touchdown site (A104-009008) contains anhydrous primary minerals of olivine and Mg-Al spinel. The other minerals included in the piece are mainly iron sulfides, magnetite, dolomite, Ca-phosphate, and ilmenite. Figure S8 shows transmission electron microscopy (TEM) images of an MgO-rich olivine crystal (Mg# of ~100) found at the surface of the Ryugu piece. The olivine grain has a nearly euhedral shape with smooth straight crystal surfaces. Most of the boundaries between the olivine crystal and the surrounding phyllosilicate-rich matrix are sharp (fig. S8C), suggesting that those olivine surfaces were not affected by aqueous alteration. We observe, however, that a small part of the olivine surface is not sharp and mixed with phyllosilicate matrix (fig. S8D). This feature may correspond to surface alteration by aqueous fluid. The presence of olivine grain with minor alteration to phyllosilicate indicates that the olivine grains accreted onto the Ryugu parent body before aqueous alteration, which took place ~5 Ma after solar system formation (3). These observations further indicate that the survived olivine grains in Ryugu experienced very minor alteration to phyllosilicates.

On an oxygen three-isotope diagram, δ^{17} O versus δ^{18} O, the compositions of primary minerals in Ryugu and Ivuna are distributed along slope-1 line (Fig. 3). Most data plot closer to the primitive chondrule mineral line (13) rather than to the carbonaceous chondrite anhydrous mineral line (11). The Δ^{17} O values range from ~-24 to ~-2% for Ryugu and from ~-24 to ~0% for Ivuna. Among the Ryugu monomineralic grains, the Δ^{17} O values

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Fig. 1. Histograms of Δ^{17} O for olivine grains. Chondrule olivine in (A) CV chondrite Kaba [data from (16)], (B) CM chondrite Murchison [data from (15)], (C) CO chondrite Yamato 81020 [data from (14)], and (D) ungrouped carbonaceous chondrite Acfer 094 [data from (13)]. Isolated olivine grains in (E) CI chondrites Ivuna and Orgueil [previous studies: data from (4, 6)], (F) CI chondrites Ivuna and Alais [previous study: data from (7)], (G) Ryugu [previous study: data from (8)], (H) Ryugu (this study), (I) Ivuna (this study), (J) comet 81P/Wild2 [data from (42, 56–60)], and (K) ungrouped carbonaceous chondrite Tagish Lake [data from (49)]. Magnesium-rich olivine (Mg# > 97) is shown as orange and Mg-poor olivine (Mg# < 97) in blue, except for (F) and (G) because their chemical compositions are not fully available. Bin sizes of (E) and (J) correspond to their analytical uncertainties. Note that Mg-poor olivine peaks in Yamato 81020 (C) and Acfer 094 (D) may be enhanced because these studies selectively measured Fe-rich ones from the polished sections. The olivine grains with low Δ^{17} O are most likely related to refractory inclusions (CAls and AOAs), while those with high Δ^{17} O are related to chondrules. L1997, Leshin *et al.* (4); P2020, Piralla *et al.* (6); M2022, Morin *et al.* (7); N2022, Nakamura *et al.* (8).



of olivine show a bimodal distribution (Fig. 1H): Three grains are $^{16}\text{O-poor}$ ($\Delta^{17}\text{O}$ of $\sim\!-5\%$), and three are $^{16}\text{O-rich}$ ($\Delta^{17}\text{O}$ of $\sim\!-23\%$). Both $^{16}\text{O-poor}$ and $^{16}\text{O-rich}$ grains coexist in the same clasts (Fig. 2). The low-Ca pyroxene is $^{16}\text{O-poor}$ ($\Delta^{17}\text{O}$ of $\sim\!-4\%$). The Mg-Al spinel is $^{16}\text{O-rich}$ ($\Delta^{17}\text{O}$ of $\sim\!-23\%$), whereas the Cr-spinel is $^{16}\text{O-poor}$ ($\Delta^{17}\text{O}$ of $\sim\!-2\%$). The bimodal distribution of $\Delta^{17}\text{O}$ is also observed among the Ivuna olivine grains

(Fig. 1I): Twenty-one grains are $^{16}\text{O-poor}$ ($\Delta^{17}\text{O}$ of \sim -7 to \sim 0%), and 7 are $^{16}\text{O-rich}$ ($\Delta^{17}\text{O}$ of \sim -23%o). The two low-Ca pyroxene grains measured in Ivuna are $^{16}\text{O-poor}$ ($\Delta^{17}\text{O}$ of \sim -5%o). The Mg-Al spinel is $^{16}\text{O-rich}$ ($\Delta^{17}\text{O}$ of \sim -23%o). The $^{16}\text{O-rich}$ olivine grains in Ryugu and Ivuna have Mg# > 97, whereas the $^{16}\text{O-poor}$ olivine grains show much larger range of Mg#, from \sim 57 to \sim 99 (fig. S10).

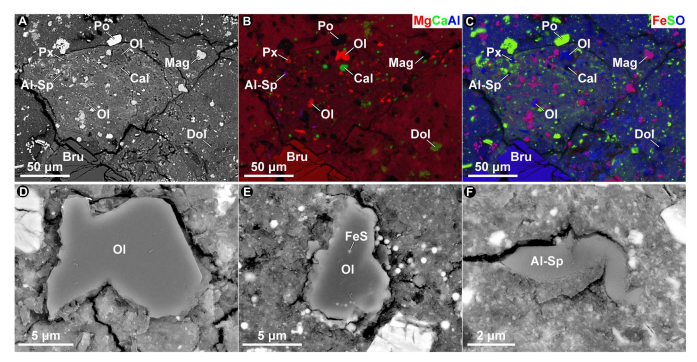


Fig. 2. Occurrences of anhydrous primary minerals in Ryugu sample. (**A**) BSE image of primary mineral-rich clast. (**B**) Combined x-ray elemental map of (A) using Mg Kα, Ca Kα, and Al Kα lines assigned for RGB color channels. (**C**) Combined x-ray elemental map of (A) using Fe Kα, S Kα, and O Kα lines assigned for RGB color channels. BSE images of (**D**) and (**E**) olivine and (**F**) Mg-Al spinel. The olivine grains (D and E) are located in the clast shown in (A) to (C). Their O isotopic compositions (Δ^{17} O) are (D) -24%, (E) -4%, and (F) -23%, respectively. Al-Sp, Mg-Al spinel; Bru, breunnerite; Cal, calcite; Dol, dolomite; FeS, Fe-sulfide; Mag, magnetite; Ol, olivine; Po, pyrrhotite; Px, low-Ca pyroxene.

A subrounded AOA, composed of nearly pure forsterite (Mg# of ~99), diopside, anorthite, and tiny Mg-Al spinel grains, was also found in Ivuna (Fig. 4, A and C). This is the first discovery of an AOA in CI chondrites. The texture and mineralogy of the AOA is like those in primitive carbonaceous chondrites (10). We also found an inclusion composed of forsteritic olivine (Mg# > 98), Mg-Al spinel, and interstitial phyllosilicates in Ivuna (Fig. 4, B and D). The texture and mineralogy of this inclusion resemble AOAs enclosing small spinel-anorthite \pm melilite-diopside—bearing CAIs (10), which experienced extensive aqueous alteration resulting in replacement of CAI anhydrous silicates by phyllosilicates. The O isotopic compositions of the Ivuna AOA and spinel-olivine inclusion plot along ~slope-1 line and have nearly identical Δ^{17} O, $-23.6 \pm 0.3\%$ (2σ , n = 5) and $-24.0 \pm 1.7\%$ (n = 3), respectively (Fig. 4, E and F).

DISCUSSION

Bimodal distributions of O isotopic compositions of primary minerals

The bimodal distribution of Δ^{17} O for the Ivuna olivines determined here based on in situ analyses on a polished section and the peak Δ^{17} O value of \sim -6‰ among 16 O-poor olivines (Fig. 1I) contrasts with previously reported data for olivine grains (>50 µm, n=19) isolated from the CI chondrites Orgueil and Ivuna (4, 6), where only 16 O-poor olivines (Δ^{17} O range from \sim -6 to +3‰ with a peak value of \sim 0‰) were observed (Fig. 1E). The observed differences between these datasets may reflect a sample bias. Comprehensive mineralogical studies of olivine grains in CI chondrites Alais,

Orgueil, and Ivuna show that most olivine grains are <30 μ m in size (5, 18, 19). Similarly, all olivine grains observed in the polished sample of Ivuna in this study are <30 μ m. Seventy-nine percent of olivines in this study have Mg# > 97 (fig. S9B), while 77% of olivines in (5, 18, 19) have Mg# > 97 (n = 69) (fig. S9C). These consistencies of size and chemical distribution indicate that our dataset of olivine grains from Ivuna is representative of CI chondrites. In contrast, the CI olivine grains measured for O isotopic compositions previously were hand-picked from disaggregated meteorites, were larger than 50 μ m, and had predominantly Mg# < 97 (Fig. 1E and fig. S9D) (4, 6). Therefore, literature data (4, 6) may represent biased sampling and are probably not representative for CI chondrites. A recent study (7) also found that both 16 O-poor and 16 O-rich olivines from the CI chondrites Ivuna and Alais by in situ O isotope analysis of those grains embedded in the matrices (Fig. 1F).

Liu *et al.* (9) argued that O isotopic exchange might have occurred in olivine grains in Ryugu. Similarly, Mg-Fe exchange during aqueous alteration (3) might have caused an Fe-enrichment in the grains. Such exchanges are, however, unlikely due to the slow volume diffusivity of cations and oxygen in olivine at the temperature relevant to aqueous alteration in Ryugu. Using the Mg-Fe interdiffusion coefficient for olivine (20), Mg-Fe exchange at a temperature of ~300°C, which is a much higher temperature than the inferred maximum temperatures of aqueous alteration on the CI parent body and Ryugu (<~150°C) (2, 4, 21), is calculated to occur in a 1-µm scale in olivine for 10¹⁰ years, which is unreasonably long for the aqueous alteration. Moreover, some olivine grains in Ryugu and Ivuna are chemically zoned, but this cannot be explained by diffusion. The Mg-Fe chemical zonings are often observed in

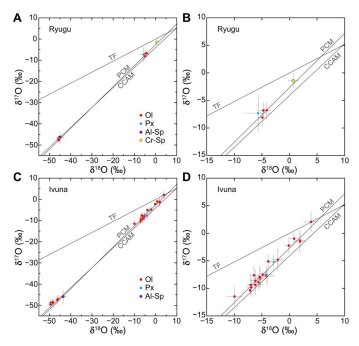


Fig. 3. Oxygen isotopic compositions of anhydrous primary minerals. (A and B) Ryugu. (C and D) Ivuna. Data are listed in tables S2 and S3 and data S1. Duplicate analyses for each grain showed identical value (within uncertainty of our measurements), suggesting homogeneous O isotopic compositions within grain. Therefore, each point corresponds to a single grain. Errors correspond to 2σ . TF, terrestrial fractionation line; CCAM, carbonaceous chondrite anhydrous mineral line; PCM, primitive chondrule mineral line.

olivine grains in ferroan porphyritic (type II) chondrules in primitive carbonaceous chondrites (13, 14, 22). The microscale heterogeneity of Mg-Fe distributions in olivine grains (fig. S6, C and D) indicates that the diffusion distance of Mg-Fe in olivine was less than a 1-µm scale. Therefore, effect of the Mg-Fe exchange is negligible in olivine. Oxygen self-diffusion rates are much smaller than the Mg-Fe interdiffusion rates in olivine for all temperatures (23). Oxygen isotopic exchange is, therefore, also excluded in olivine. Similarly, effects of diffusive exchanges of Mg-Fe and O isotopes in pyroxene and spinel by solid-state diffusion are negligible at the temperatures experienced by the CI parent body and Ryugu (24–27). Therefore, the chemical and O isotopic compositions of olivine, low-Ca pyroxene, and spinel in Ryugu and Ivuna have not been disturbed after accretion, and they most likely preserve preaccretion signatures acquired in the solar nebula environment.

To summarize, olivine grains in Ryugu show a bimodal distribution of Δ^{17} O, \sim 23 and \sim 5‰ (Fig. 1H), similar to that observed for olivine grains in Ivuna (Fig. 1I). The O isotopic compositions of low-Ca pyroxene and Mg-Al spinel in Ryugu are also similar to those measured in Ivuna minerals (Fig. 3). We conclude that primary minerals in Ryugu and CI chondrites have similar characteristics and sampled two isotopically distinct materials, $^{16}\text{O}\text{-rich}$ and $^{16}\text{O}\text{-poor}.$

Origin of ¹⁶O-poor primary minerals in Ryugu and Ivuna

The ^{16}O -poor olivine and low-Ca pyroxene in Ryugu and Ivuna have $\Delta^{17}\text{O}$ values ranging from \sim -7 to \sim 0% (Fig. 1, H and I). This range is consistent with those observed for olivine and low-

Ca pyroxene in porphyritic chondrules from several carbonaceous chondrite groups [CV (Vigarano-type), CM (Mighei-type), and CO (Ornans-type)] and the ungrouped carbonaceous chondrite Acfer 094 (Fig. 1, A to D). The Ivuna olivine grains with Mg# < 97 have typically higher $\Delta^{17}O$ (>~-2‰) than more Mg-rich olivine (~-6%) (Fig. 1I and fig. S10). Such relationships between Δ^{17} O and Mg# are often observed for olivine (Fig. 1, A to D) and low-Ca pyroxene phenocrysts in chondrules from carbonaceous chondrites (13–17). Moreover, the most frequent Δ^{17} O value (~-6‰) for the Ivuna olivine is identical to those for chondrules (Fig. 1, A to D and I). The chemical features of the Ryugu and Ivuna olivine grains are consistent with those observed for olivine in chondrules of the primitive carbonaceous chondrites (fig. S11). Fe,Ni metal blebs in forsteritic grain seen in the Ivuna sample (fig. S7A) are frequently observed in Mg-rich porphyritic chondrules from carbonaceous chondrites (13–17). The Ryugu Cr-spinel has a Δ^{17} O value of $-1.8 \pm 1.4\%$. Chrome-spinel is commonly observed in type II chondrules in carbonaceous chondrites (28). The majority of type II chondrules in carbonaceous chondrites have a Δ^{17} O of ~-2‰ (13-17). Therefore, we infer that the Ryugu Cr-spinel most likely originated from a carbonaceous chondrite type II chondrule-like melt. Similar to CIs, Ryugu is devoid of chondrules, but the ¹⁶Opoor olivine, low-Ca pyroxene, and Cr-spinel grains in Ryugu

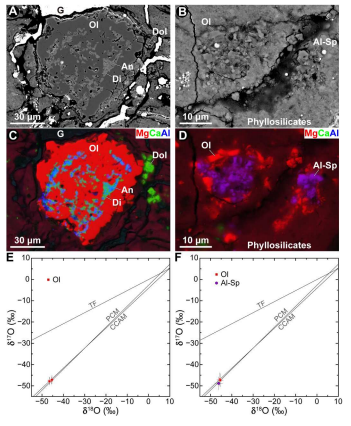


Fig. 4. Refractory inclusions in Ivuna. (**A** and **B**) BSE images and (**C** and **D**) combined x-ray elemental maps of using Mg K α , Ca K α , and Al K α lines assigned for RGB color channels of (A and C) AOA and (B and D) spinel-olivine inclusion from Ivuna. Oxygen isotopic compositions of individual minerals in (**E**) the AOA and (**F**) the spinel-olivine inclusion. Errors correspond to 2σ . An, anorthite; Di, diopside; Dol, dolomite; Ol, olivine; G, gold coating residue.

and Ivuna most likely represent fragments of chondrule-like objects.

Origin of ¹⁶O-rich primary minerals in Ryugu and Ivuna

The AOA and spinel-olivine inclusion found in Ivuna (Fig. 4) are textually, mineralogically, and isotopically similar to AOAs and spinel-rich, fine-grained CAIs in carbonaceous chondrites (10, 29-33). These objects most likely formed by condensation from an ¹⁶O-rich gaseous reservoir. The presence of interstitial phyllosilicates in the spinel-olivine inclusion suggests that it subsequently experienced aqueous alteration, probably on the Ivuna parent body. The ¹⁶O-rich isolated olivine and Mg-Al spinel grains in Ryugu and Ivuna studied here have Δ^{17} O values identical to those of the Ivuna AOA-like objects and to those of most CAIs and AOAs from other types of carbonaceous chondrites (31, 33–35); they consistently show Δ^{17} O of ~-23 to -24‰. Ryugu ¹⁶O-rich olivine grains presented by Nakamura et al. (8) also show Δ^{17} O of ~-23‰ (Fig. 1G). In contrast, ¹⁶O-rich olivine and low-Ca pyroxene in CI chondrites by Morin et al. (7) show variations in Δ^{17} O ranging from -24 to -19% (Fig. 1F). The variations are clearly larger than their given analytical uncertainties of typically ~0.6%. Further studies are needed to clarify the origin or reason of variable Δ^{17} O for ¹⁶O-rich minerals.

The ¹⁶O-rich isolated olivine grains studied here are all Mg-rich (Mg# > 97) (fig. S10), and their CaO contents are <~0.15 weight % (fig. S11), consistent with compositions for olivine grains of the Ivuna AOA-like objects and those of AOAs from primitive carbonaceous chondrites (10, 30, 31). Irregular shapes of Mg-Al spinel grains in Ryugu and Ivuna (figs. S6F and S7C) are consistent with mineralogical textures of spinel-rich, fine-grained CAIs in carbonaceous chondrites (29, 31, 32). These observations indicate that the ¹⁶O-rich olivine and Mg-Al spinel grains in Ryugu and Ivuna represent fragments of refractory inclusions, and/or isolated grains condensed in the refractory inclusion-forming region. Moreover, the CAI-like mineral assemblages composed of Mg-Al spinel, perovskite, and ±hibonite were reported in the C0040 and C0002 Ryugu samples (2). A fragment of melilite-rich (type A) CAI was found in Ivuna (36). Micrometer-sized ¹⁶O-rich (Δ¹⁷O of ~-23‰) corundum grains were reported in acid residues of Orgueil (37). The presence of ¹⁶O-rich refractory inclusion-like objects and isolated mineral grains in Ryugu and Ivuna indicates that fragments of ¹⁶O-rich refractory inclusions were also clearly one of the building blocks of the Ryugu and Ivuna parent body. The presence of both ¹⁶O-poor chondrule-like and ¹⁶O-rich refractory inclusion-like minerals in Ryugu and Ivuna suggests that some of building blocks of Ryugu and Ivuna are similar to those of other carbonaceous chondrite groups.

Implications for the accretion region of Ryugu and Ivuna parent bodies

The ¹⁶O-rich objects in Ryugu and Ivuna, texturally and isotopically similar to refractory inclusions, most likely formed in a high-temperature, innermost region of the solar protoplanetary disk, possibly <~0.1 au (astronomical units) from the proto-Sun (38) where the ambient gas was primarily ¹⁶O-rich (12). The accretion regions of the Ryugu and CI chondrites parent body are inferred to be beyond 3 to 4 au (2) and possibly as far as ~15 au from the Sun (39). Whole rock Fe isotopic compositions of Ryugu and CI chondrites also imply that they are accreted in the outer Solar System (40). In this

context, the ¹⁶O-rich primary minerals were transported outward from the innermost region to the outerskirts of the disk system. The similar conclusion was reached to explain the presence of high-temperature minerals in the comet 81P/Wild2 samples (41, 42).

Normal-sized chondrules (~100 to ~2000 µm in apparent diameter) (43) that are commonly observed in most groups of carbonaceous chondrites, as well as their pseudomorphs, have not been identified in Ryugu and CI chondrites [(2-9); this study]. This cannot be explained by extensive aqueous alteration experienced by the Ryugu and CI parent bodies, because chondrule pseudomorphs are preserved in the nearly completely aqueously altered CM and CR (Renazzo-type) carbonaceous chondrites (44, 45). Furthermore, isotopic fractionations of moderately volatile elements in CIs are consistent with accretion from material largely devoid of chondrules (46). The ¹⁶O-poor olivine, low-Ca pyroxene, and Crspinel in Ryugu and Ivuna are typically <30 μm in size and most likely represent chondrule fragments. These observations suggest that chondrule-forming events were rare at the accretion time and/or near the accretion region of the Ryugu and CI parent bodies compared to those of other carbonaceous chondrite groups.

The ratios of the refractory inclusion–like ¹⁶O-rich olivine grains to the chondrule-like ¹⁶O-poor olivine grains studied here are 3:3 for Ryugu and 7:21 for Ivuna, respectively (Fig. 1, H and I). If we combined with the data of Nakamura et al. (8)(Fig. 1G), the proportion for Ryugu is 6:7. These proportions are like that for olivine grains of the comet 81P/Wild2 samples of 4:10 (Fig. 1J). On the other hand, the ratios for Ryugu, Ivuna, and comet 81P/Wild2 are in sharp contrast with ratios for other carbonaceous chondrite groups. In the highly hydrated carbonaceous chondrite Tagish Lake (C-ungrouped) that contains rare chondrules (47) and was inferred to have accreted at >10 au from the Sun (48), the ratio for isolated olivine grains, embedded in matrices, is 1:21 (Fig. 1K) (49); ¹⁶O-rich olivine is much rarer than those for Ryugu, Ivuna, and comet 81P/Wild2. In other carbonaceous chondrites (CO, CV, CM, and C-ungrouped), none of 31 isolated olivine grains exhibit refractory inclusion-like ¹⁶O-rich compositions; instead, all of them show chondrule-like ¹⁶O-poor compositions (50). If we estimate the abundance of refractory inclusion-like ¹⁶O-rich olivine among all olivine grains, that for Ryugu and Ivuna is $32 \pm 15\%$ (2 σ). The abundance for other carbonaceous chondrite groups (CO, CV, CM, and C-ungrouped including Tagish Lake) is 2 \pm 4% and that for 81P/Wild2 samples is 29 \pm 24%. The abundance of refractory inclusion-like olivine for Ryugu and Ivuna and that for 81P/Wild2 samples are very similar to each other. On the other hand, that for Ryugu and Ivuna is clearly different from that for other carbonaceous chondrite groups. Moreover, in CV carbonaceous chondrites, characterized by the highest abundance of refractory inclusions among carbonaceous chondrites, the volume ratio of refractory inclusions to chondrules is 1:10 to 1:5 (51, 52).

The ratio for olivine in Ryugu and Ivuna that can be tied to refractory inclusions and chondrules is like what has been documented in comet 81P/Wild2. The relative paucity of chondrule-derived anhydrous grains in Ryugu, Ivuna, and 81P/Wild2 could be due to efficient outward transport of refractory inclusion—derived grains (37) formed at the earliest stage of evolution of the solar protoplanetary disk (31, 32) and/or rarer chondrule-formation events near the accretion regions. We infer that the accretion region of Ryugu and CI parent bodies is distinct from those for other carbonaceous

chondrite groups including Tagish Lake and could be closer to the accretion region of 81P/Wild2 comet.

MATERIALS AND METHODS

Sample preparation

Polished sections of C0002-C1001 (fig. S3) and Ivuna-HK2 (fig. S5) made from the Ryugu sample C0002 and the Ivuna CI chondrite were used for mineralogical and petrological observations and in situ O isotope measurements by secondary ion mass spectrometry (SIMS). The samples were individually embedded in 25mm epoxy disks using the Buehler EpoxiCure 2 resin. After embedding, their sample surface sides were also impregnated with the resin in vacuum, to avoid collapsing the fragile samples during polishing. The sample disks were polished with an automatic polishing machine (Musashino Denshi MA-200e) at Hokkaido University. Diamond slurry with polycrystalline diamond particles of ~3 μm dissolved in ethylene glycol sprayed on a copper polishing plate was used to obtain flat surface of the sample disks. During the flattening, the sample surfaces were impregnated with the resin in vacuum a few times. Subsequently, ~1-μm diamond slurry sprayed on a tin-antimony alloy polishing plate and on a polishing cloth was used to finalize the polishing. These procedures were adopted to obtain flat and smooth surface for anhydrous minerals, which is critical for the quality of SIMS measurements. Only >99.5% ethanol was used for cleaning during and after the polishing. The polished sections were coated with a thin (~5 nm) gold film using a Leica EM ACE600 coater at Hokkaido University, for backscattered electron (BSE) and x-ray imaging, and elemental analysis before in situ O isotope measurements.

Electron microscopy

BSE images were obtained using a field-emission scanning electron microscope (FE-SEM; JEOL JSM-7000F) at Hokkaido University. X-ray elemental analyses were conducted with a 15-keV electron beam using an energy dispersive x-ray spectrometer (EDS; Oxford X-Max 150) installed on the FE-SEM. Beam currents of ~2 and ~1 nA were used for x-ray mapping and quantitative analysis, respectively. Quantitative calculations were conducted using Oxford AZtec software. We used in-house standards, such as San Carlos olivine and Mn metal, for the standardization. X-ray elemental maps covering the entire polished sections of C0002-C1001 (fig. S3) and Ivuna-HK2 (fig. S5) were obtained with pixel sizes of 0.24 and 0.48 µm, respectively, to systematically find out olivine, pyroxene, and spinel grains that can be measured for O isotopic compositions with SIMS. Representative chemical compositions of the primary minerals are shown in table S1. After electron microscopy was completed, the polished sections were recoated with an additional thin (~65 nm) gold film for SIMS measurements.

For the TEM of olivine in Ryugu, we used a fine grain (A104-009008) from the first touchdown site. We observed the grain using FE-SEM (JEOL JSM-7001F) at Kyoto University. Then, electron-transparent sections were extracted from the region of interest on the Ryugu piece using a focused ion beam (FIB) system (Helios NanoLab G3 CX at Kyoto University). The Ryugu piece was coated with an electron-beam–deposited Pt layer (at 2 kV), followed by a Ga ion-beam–deposited Pt layer (at 30 kV). The sections of a few tens of micrometers in size were extracted and were thinned to 50 to 200 nm using a 16- to 30-keV Ga⁺ beam and lastly cleaned using a

2-keV Ga^+ beam at 77 pA. The FIB sections were analyzed using an FE-TEM (JEOL JEM 2100F) equipped with an EDS (JEOL JED-2300 T) at Kyoto University. Bright-field TEM images, selected-area electron diffraction patterns, and scanning transmission electron microscopy (STEM) images at 200 kV were obtained using a charge-coupled device or complementary metal oxide semiconductor camera (Gatan Orius200D, Rio9). EDS analysis was performed in STEM mode. Quantitative elemental abundances were calculated using the ζ -factor method (53).

In situ O isotope measurements by SIMS

The O isotopic compositions of olivine, pyroxene, and spinel in Ryugu and Ivuna were measured in situ with the Cameca ims-1280HR SIMS instrument at Hokkaido University. The analytical and instrumental settings were established by Kawasaki *et al.* (35). The polished sections were coated with a thin (~70 nm) gold film. Measurement spots are shown in figs. S12 to S51. A $^{133}\mathrm{Cs^+}$ primary beam accelerated to 20 keV was used in the experiment. A normal-incidence electron flood gun was used for electrostatic charge compensation of the analyzing areas during the measurements. Negative secondary ions ($^{16}\mathrm{O}^-$, $^{17}\mathrm{O}^-$, and $^{18}\mathrm{O}^-$) were measured simultaneously in the multicollection mode. The mass resolution of $M/\Delta M$ for $^{17}\mathrm{O}^-$ was set at >6000 to resolve $^{17}\mathrm{O}^-$ from $^{16}\mathrm{OH}^-$ while that for $^{16}\mathrm{O}^-$ and $^{18}\mathrm{O}^-$ was ~2000. The automatic centering program (dynamic transfer deflector and magnetic field) was applied before data collection.

Analyzed areas were precisely determined according to scanning ion image of $^{16}\mathrm{O}^-$ collected by a multicollector electron multiplier (EM; designated as L2), which was not used for the data collection. Before measurements, we made a few sputtered craters near measurement targets using a 10- to 30-pA primary beam by the SIMS, and then electron images were obtained by the FE-SEM to obtain distances from the sputtered craters to the measurement targets. The craters were visible in $^{16}\mathrm{O}^-$ scanning images and used to locate the target minerals.

The reported uncertainties in the O-isotopic compositions were the larger of the external reproducibility of standard measurements (2 SD) or internal precision (2 SE of cycle data) of samples. Measurement spots were observed by the FE-SEM after SIMS measurements. The data from spots with inclusions and overlapping matrix minerals were rejected. The reasons of all the rejected spots are written in figs. S12 to S51 and data S1.

We used three conditions with different primary beam currents depending on mineral sizes. An ~1-nA primary beam with an elliptical shape of 6 µm by 8 µm was used for the measurement of olivine grains in Ivuna. The primary beam was rastered over an 8 µm by 8 μm area during the presputtering for 60 s, and then the raster size was reduced to 1 μm by 1 μm for the data collection. $^{16}O^{-}$, ¹⁷O⁻, and ¹⁸O⁻ were measured using a multicollector Faraday cup (FC; 10¹⁰ ohm, designated as L'2), an axial FC (10¹² ohm), and a multicollector FC (1012 ohm, designated as H1), respectively. The secondary ion intensity of $^{16}O^-$ was ~1.5 to 1.7×10^9 counts per second (cps). The data were collected for 30 cycles with 4-s integration time per cycle. Obtained count rates were corrected for the FC background, monitored during the presputtering of every measurement, and relative yield of each detector. The ¹⁶OH⁻ count rate was measured immediately after the measurements, and we made a small tail correction on ¹⁷O⁻, although its contribution to ¹⁷O⁻

was smaller than ~0.1‰. Typical uncertainties for δ^{17} O, δ^{18} O, and Δ^{17} O were 0.6, 0.3, and 0.6‰ (2 σ), respectively.

An ~30-pA primary beam with an elliptical shape of ~2 µm by 3 µm (~2.5 µm by 3.5 µm including the beam halo) was used for the measurement of olivine, pyroxene, and Cr-spinel in Ryugu and Ivuna. $^{16}\mathrm{O}^-$, $^{17}\mathrm{O}^-$, and $^{18}\mathrm{O}^-$ were measured using a multicollector FC (10¹¹ ohm, designated as L1), an axial EM, and a multicollector EM (designated as H2), respectively. The secondary ion intensities of $^{16}\mathrm{O}^-$ were ~1.7 to 2.9 × 10⁷, ~2.2 × 10⁷, and ~ 2.9 × 10⁷ cps for olivine, pyroxene, and Cr-spinel, respectively. The data were collected for 60 cycles with 4-s integration time per cycle. Obtained count rates were corrected for the FC background, EM dead time, and relative yield of each detector. Typical uncertainties for $\delta^{17}\mathrm{O}$, $\delta^{18}\mathrm{O}$, and $\Delta^{17}\mathrm{O}$ were 1.6, 1.0, and 1.7%, respectively.

An ~3-pA primary beam with an elliptical shape of 0.8 μm by 1.3 μm (~1.0 μm by 2.0 μm including the beam halo) was used for the measurement of olivine, pyroxene, and Mg-Al spinel in Ryugu and Ivuna. Detector settings are the same with those for the ~30-pA condition above. The secondary ion intensities of $^{16}O^-$ were ~1.5 to 2.1 \times 10⁶ cps. The data were collected for 200 cycles with 4-s integration time per cycle. The $^{16}OH^-$ count rate was measured immediately after the measurements, but we did not make a tail correction on $^{17}O^-$ because its contribution to $^{17}O^-$ was calculated as typically ~0.03%o and up to ~0.2%o, comparable to that for standards. Typical uncertainties for $\delta^{17}O$, $\delta^{18}O$, and $\Delta^{17}O$ were 3.1, 2.0, and 3.1%o, respectively.

San Carlos olivine (Mg# = 89; $\delta^{18}O = 5.2\%$), synthetic enstatite ($\delta^{18}O = 10.55\%$), and Russian spinel ($\delta^{18}O = 8.5\%$) (54) were used as standards to correct the instrumental mass fractionation for olivine, pyroxene, and spinel, respectively. Because Mg# of olivine grains are larger than 76, except for an olivine grain in Ivuna with Mg# of ~57, variations in instrumental mass fractionation correlated with Mg# of olivine from that of San Carlos olivine (55) could be insignificant in the analytical uncertainties of this study.

Supplementary Materials

This PDF file includes:

Figs. S1 to S51 Tables S1 to S3

Other Supplementary Material for this manuscript includes the following:
Data S1

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