

Title	Carbonate record of temporal change in oxygen fugacity and gaseous species in asteroid Ryugu
Author(s)	Fujiya, Wataru; Kawasaki, Noriyuki; Nagashima, Kazuhide; Sakamoto, Naoya; O'D. Alexander, Conel M.; Kita, Noriko T.; Kitajima, Kouki; Abe, Yoshinari; Aleon, Jerome; Amari, Sachiko; Amelin, Yuri; Bajo, Ken-ichi; Bizzarro, Martin; Bouvier, Audrey; Carlson, Richard W.; Chaussidon, Marc; Choi, Byeon-Gak; Dauphas, Nicolas; Davis, Andrew M.; Di Rocco, Tommaso; Fukai, Ryota; Gautam, Ikshu; Haba, Makiko K.; Hibiya, Yuki; Hidaka, Hiroshi; Homma, Hisashi; Hoppe, Peter; Huss, Gary R.; Ichida, Kiyohiro; Iizuka, Tsuyoshi; Ireland, Trevor R.; Ishikawa, Akira; Itoh, Shoichi; Kleine, Thorsten; Komatani, Shintaro; Krot, Alexander N.; Liu, Ming-Chang; Masuda, Yuki; McKeegan, Kevin D.; Morita, Mayu; Motomura, Kazuko; Moynier, Frederic; Nakai, Izumi; Nguyen, Ann; Nittler, Larry; Onose, Morihiko; Pack, Andreas; Park, Changkur; Piani, Laurette; Qin, Liping; Russell, Sara S.; Schonbachler, Maria; Tafla, Lauren; Tang, Haolan; Terada, Kentaro; Terada, Yasuko; Usui, Tomohiro; Wada, Sohei; Wadhwa, Meenakshi; Walker, Richard J.; Yamashita, Katsuyuki; Yin, Qing-Zhu; Yokoyama, Tetsuya; Yoneda, Shigekazu; Young, Edward D.; Yui, Hiroharu; Zhang, Ai-Cheng; Nakamura, Tomoki; Naraoka, Hiroshi; Noguchi, Takaaki; Okazaki, Ryuji; Sakamoto, Kanako; Yabuta, Hikaru; Abe, Masanao; Miyazaki, Akiko; Nakato, Aiko; Nishimura, Masahiro; Okada, Tatsuaki; Yada, Toru; Yogata, Kasumi; Nakazawa, Satoru; Saiki, Takanao; Tanaka, Satoshi; Terui, Fuyuto; Tsuda, Yuichi; Watanabe, Sei-ichiro; Yoshikawa, Makoto; Tachibana, Shogo; Yurimoto, Hisayoshi
Citation	Nature Geoscience, 16(8), 675-682 https://doi.org/10.1038/s41561-023-01226-y
Issue Date	2023-08
Doc URL	http://hdl.handle.net/2115/91054
Туре	article (author version)
File Information	Fujiya_Ryugu_carbonates_rev2_final.pdf



Carbonate record of temporal change in oxygen fugacity and gaseous species in asteroid Ryugu

3

4 Wataru Fujiya^{1*}, Noriyuki Kawasaki², Kazuhide Nagashima³, Naoya Sakamoto⁴, Conel M. O'D. 5 Alexander⁵, Noriko T. Kita⁶, Kouki Kitajima⁶, Yoshinari Abe⁷, Jérôme Aléon⁸, Sachiko Amari^{9,10}, Yuri Amelin¹¹, Ken-ichi Bajo², Martin Bizzarro¹², Audrey Bouvier¹³, Richard W. Carlson⁵, Marc 6 Chaussidon¹⁴, Byeon-Gak Choi¹⁵, Nicolas Dauphas¹⁶, Andrew M. Davis¹⁶, Tommaso Di Rocco¹⁷, 7 Ryota Fukai¹⁸, Ikshu Gautam¹⁹, Makiko K. Haba¹⁹, Yuki Hibiya²⁰, Hiroshi Hidaka²¹, Hisashi Homma²², 8 9 Peter Hoppe²³, Gary R. Huss³, Kiyohiro Ichida²⁴, Tsuyoshi Iizuka²⁵, Trevor R. Ireland²⁶, Akira Ishikawa¹⁹, Shoichi Itoh²⁷, Thorsten Kleine²⁸, Shintaro Komatani²⁴, Alexander N. Krot³, Ming-Chang 10 Liu^{29,30}, Yuki Masuda¹⁹, Kevin D. McKeegan²⁹, Mayu Morita²⁴, Kazuko Motomura³¹, Frédéric 11 Moynier¹⁴, Izumi Nakai³², Ann Nguyen³³, Larry Nittler⁵, Morihiko Onose²⁴, Andreas Pack¹⁷, 12 Changkun Park³⁴, Laurette Piani³⁵, Liping Qin³⁶, Sara S. Russell³⁷, Maria Schönbächler³⁸, Lauren 13 Tafla²⁹, Haolan Tang²⁹, Kentaro Terada³⁹, Yasuko Terada⁴⁰, Tomohiro Usui¹⁸, Sohei Wada², Meenakshi 14 Wadhwa⁴¹, Richard J. Walker⁴², Katsuyuki Yamashita⁴³, Qing-Zhu Yin⁴⁴, Tetsuya Yokoyama¹⁹, 15 Shigekazu Yoneda⁴⁵, Edward D. Young²⁹, Hiroharu Yui⁴⁶, Ai-Cheng Zhang⁴⁷, Tomoki Nakamura⁴⁸, 16 Hiroshi Naraoka⁴⁹, Takaaki Noguchi²⁷, Ryuji Okazaki⁴⁹, Kanako Sakamoto¹⁸, Hikaru Yabuta⁵⁰, 17 Masanao Abe¹⁸, Akiko Miyazaki¹⁸, Aiko Nakato¹⁸, Masahiro Nishimura¹⁸, Tatsuaki Okada¹⁸, Toru 18 Yada¹⁸, Kasumi Yogata¹⁸, Satoru Nakazawa¹⁸, Takanao Saiki¹⁸, Satoshi Tanaka¹⁸, Fuyuto Terui⁵¹, 19 20 Yuichi Tsuda¹⁸, Sei-ichiro Watanabe²¹, Makoto Yoshikawa¹⁸, Shogo Tachibana⁵², and Hisayoshi 21 Yurimoto²

22

23 *Corresponding author (wataru.fujiya.sci@vc.ibaraki.ac.jp)

24

25 Affiliations:

- ²⁶ ¹Faculty of Science, Ibaraki University; Mito 310-8512, Japan.
- ²Department of Natural History Sciences, Hokkaido University; Sapporo 060-0810, Japan.
- ³Hawai'i Institute of Geophysics and Planetology, University of Hawai'i at Mānoa; Honolulu, HI
 96822, USA.
- ⁴Isotope Imaging Laboratory, Creative Research Institution, Hokkaido University; Sapporo 001-0021,
 Japan.
- ⁵Earth and Planets Laboratory, Carnegie Institution for Science; Washington, DC, 20015, USA.
- ³³ ⁶Geoscience, University of Wisconsin- Madison; Madison, WI 53706, USA.
- ⁷Graduate School of Engineering Materials Science and Engineering, Tokyo Denki University; Tokyo
 120-8551, Japan.
- 36 ⁸Institut de Minéralogie, de Physique des Matériaux et de Cosmochimie, Sorbonne Université,

- 37 Museum National d'Histoire Naturelle, CNRS UMR 7590, IRD; 75005 Paris, France.
- ⁹McDonnell Center for the Space Sciences and Physics Department, Washington University; St. Louis,
 MO 63130, USA.
- 40 ¹⁰Geochemical Research Center, The University of Tokyo; Tokyo 113-0033, Japan.
- ¹¹Guangzhou Institute of Geochemistry, Chinese Academy of Sciences; Guangzhou, GD 510640,
 China.
- ¹²Centre for Star and Planet Formation, GLOBE Institute, University of Copenhagen; Copenhagen, K
 1350, Denmark.
- ⁴⁵ ¹³Bayerisches Geoinstitut, Universität Bayreuth; Bayreuth 95447, Germany.
- ⁴⁶ ¹⁴Université Paris Cité, Institut de physique du globe de Paris, CNRS; 75005 Paris, France
- ⁴⁷ ¹⁵Department of Earth Science Education, Seoul National University; Seoul 08826, Republic of Korea.
- 48 ¹⁶Department of the Geophysical Sciences and Enrico Fermi Institute, The University of Chicago,
- 49 5734 South Ellis Avenue, Chicago 60637, USA.
- ⁵⁰ ¹⁷Faculty of Geosciences and Geography, University of Göttingen; Göttingen, D-37077, Germany.
- ¹⁸ISAS/JSEC, JAXA; Sagamihara 252-5210, Japan.
- ¹⁹Department of Earth and Planetary Sciences, Tokyo Institute of Technology; Tokyo 152-8551, Japan.
- ²⁰General Systems Studies, The University of Tokyo; Tokyo 153-0041, Japan.
- ⁵⁴ ²¹Earth and Planetary Sciences, Nagoya University; Nagoya 464-8601, Japan.
- ²²Osaka Application Laboratory, SBUWDX, Rigaku Corporation; Osaka 569-1146, Japan.
- ²³Max Planck Institute for Chemistry; Mainz 55128, Germany.
- ²⁴Analytical Technology, Horiba Techno Service Co., Ltd.; Kyoto 601-8125, Japan.
- ²⁵Earth and Planetary Science, The University of Tokyo; Tokyo 113-0033, Japan.
- ²⁶School of Earth and Environmental Sciences, The University of Queensland; St Lucia QLD 4072,
 Australia.
- 61 ²⁷Earth and Planetary Sciences, Kyoto University; Kyoto 606-8502, Japan.
- ²⁸Max Planck Institute for Solar System Research; 37077 Göttingen, Germany.
- ⁶³²⁹Earth, Planetary, and Space Sciences, UCLA; Los Angeles, CA 90095, USA.
- ⁶⁴ ³⁰Lawrence Livermore National Laboratory; Livermore, CA 94550, USA.
- ⁶⁵ ³¹Thermal Analysis, Rigaku Corporation; Tokyo 196-8666, Japan.
- ³²Applied Chemistry, Tokyo University of Science; Tokyo 162-8601, Japan.
- ³³Astromaterials Research and Exploration Science, NASA Johnson Space Center; Houston, TX
 77058, USA.
- ⁶⁹ ³⁴Earth-System Sciences, Korea Polar Research Institute; Incheon 21990, Korea.
- ³⁵Centre de Recherches Pétrographiques et Géochimiques, CNRS Université de Lorraine; 54500
 Nancy, France.
- ³⁶University of Science and Technology of China, School of Earth and Space Sciences; Anhui 230026,

- 73 China.
- ³⁷Department of Earth Sciences, Natural History Museum; London, SW7 5BD, UK.
- ⁷⁵ ³⁸Institute for Geochemistry and Petrology, Department of Earth Sciences, ETH Zurich; Zurich,
- 76 Switzerland.
- ³⁹Earth and Space Science, Osaka University; Osaka 560-0043, Japan.
- ⁴⁰Spectroscopy and Imaging, Japan Synchrotron Radiation Research Institute; Hyogo 679-5198 Japan.
- ⁴¹School of Earth and Space Exploration, Arizona State University; Tempe, AZ 85281, USA.
- ⁴²Geology, University of Maryland; College Park, MD 20742, USA.
- ⁴³Graduate School of Natural Science and Technology, Okayama University; Okayama 700-8530,
 Japan.
- ⁴⁴Earth and Planetary Sciences, University of California; Davis, CA 95616, USA.
- ⁴⁵Science and Engineering, National Museum of Nature and Science; Tsukuba 305-0005, Japan.
- ⁴⁶Chemistry, Tokyo University of Science; Tokyo 162-8601, Japan.
- ⁴⁷School of Earth Sciences and Engineering, Nanjing University; Nanjing 210023, China.
- ⁴⁸Department of Earth Science, Tohoku University; Sendai, 980-8578, Japan.
- ⁴⁹Department of Earth and Planetary Sciences, Kyushu University; Fukuoka 819-0395, Japan.
- ⁵⁰Earth and Planetary Systems Science Program, Hiroshima University; Higashi-Hiroshima, 739-8526,
 Japan.
- ⁵¹Kanagawa Institute of Technology; Atsugi 243-0292, Japan.
- ⁵²UTokyo Organization for Planetary and Space Science, University of Tokyo; Tokyo 113-0033, Japan.
- 93

94 The Hayabusa2 spacecraft explored asteroid 162173 Ryugu and brought its surface materials to 95 Earth. Ryugu samples resemble Ivuna-type (CI) chondrites – the most chemically primitive 96 meteorites - and contain secondary phyllosilicates and carbonates, which are indicative of 97 aqueous alteration. Understanding the conditions (such as temperature, redox conditions, and 98 fluid composition) during aqueous alteration is crucial to elucidating how Ryugu evolved to its 99 present state, but little is known about the temporal changes in these conditions. Here we show that calcium carbonate (calcite) grains in Ryugu and Ivuna samples have variable ¹⁸O/¹⁶O and 100 ¹³C/¹²C ratios that are respectively 24–46‰ and 65–108‰ greater than terrestrial standard 101 102 values, whereas those of calcium-magnesium carbonate (dolomite) grains are much more 103 homogeneous, ranging within 31-36‰ for oxygen and 67-75‰ for carbon. We infer that the 104 calcite precipitated first over a wide range of temperatures and oxygen partial pressures, and 105 that the proportion of gaseous $CO_2/CO/CH_4$ molecules changed temporally. By contrast, the 106 dolomite formed later in a more oxygen-rich and thus CO₂-dominated environment when the 107 system was approaching equilibrium. The characteristic isotopic compositions of secondary 108 carbonates in Ryugu and Ivuna are not observed for other hydrous meteorites, suggesting a 109 unique evolutionary pathway for these asteroids.

110

111 The Japan Aerospace Exploration Agency (JAXA) Hayabusa2 spacecraft explored the near-Earth 112 asteroid 162173 Ryugu and brought samples of its surface materials back to Earth^{1,2}. Ryugu has been 113 classified spectroscopically as a member of the C-complex of asteroids^{3,4}. It is a rubble pile asteroid 114 consisting of numerous rocky blocks that are the fragments resulting from the disruption of an original, 115 larger parent body⁵⁻⁷.

116

Previous work has reported that Ryugu materials underwent extensive aqueous alteration as the result of water activity in the original parent body⁸⁻¹⁰ and are mainly composed of secondary minerals that formed during the aqueous alteration: phyllosilicates, carbonates, sulphides, and oxides. On the other hand, primary minerals like anhydrous silicates are rare⁹. These petrological characteristics are comparable to those in the CI (Ivuna-type) chondritic meteorites, pointing to a kinship between Ryugu and CI chondrites. The Ryugu's whole-rock chemical and isotope compositions confirm a close affinity with CI chondrites^{8,11,12}.

124

125 Carbonates, the major Ca budget in Ryugu and CI chondrites¹³, are of particular interest because (*i*) 126 their chemical and isotopic compositions reflect the conditions of aqueous alteration, and (*ii*) their

grain size is commonly large enough to allow *in-situ* analysis by electron and ion microprobes¹⁴⁻¹⁷.

- 128 The C source of carbonates is unclear but was likely ices that included CO, CO₂, and CH₄, and/or
- The e source of carbonates is unclear out was fixery less that included eo, eo₂, and eria, and/or
- 129 organic matter^{18,19}. These materials may have formed in the solar nebula or even the parental molecular

130 cloud of the solar system. Therefore, the C isotope compositions recorded by carbonates can help us

- 131 shed light on the physicochemical processes that operated in these environments.
- 132

133 In this study, we investigate the conditions of aqueous alteration and the origin of the materials 134 accreted by the Ryugu/CI parent bodies. To this end, we performed *in-situ* O and C isotope 135 measurements of calcite (CaCO₃) and dolomite (CaMg(CO₃)₂) in Ryugu samples A0058, collected at 136 the first touchdown site, and C0002, from the second touchdown site, as well as the Ivuna meteorite. 137 The O isotope data of the A0058 and Ivuna dolomite are taken from a previous study⁸.

138

139 Occurrences and isotope compositions of carbonate minerals

140 Dolomite is the most abundant carbonate mineral in the analysed samples, and we found numerous 141 dolomite grains throughout the Ryugu and Ivuna matrices, whereas calcite is rare. No calcite grains 142 were observed in the Ryugu A0058 sample studied. Calcite was found only in limited areas (clasts) of 143 the Ryugu C0002 and Ivuna samples, occurring with primary anhydrous silicate minerals like Mg-rich 144 olivine and pyroxene (Fig. 1a). The calcite grains (<10 µm in size) are usually smaller than the 145 dolomite grains (several tens of µm) (Fig. 1b). The dolomite has compositional variation and complex 146 zoning within grains (Extended Data Fig. 1). We also found breunnerite (Mg(Fe,Mn)(CO₃)₂) grains, 147 for which we did not measure isotope compositions because of the lack of a suitable standard material 148 for isotope analysis.

149

150 The O and C isotope compositions of the Ryugu and Ivuna carbonates are similar (Table 1). The δ^{18} O 151 values of the calcite in C0002 and Ivuna (+24 to +46‰) show a grain-to-grain variation larger than those of the dolomite in C0002, A0058, and Ivuna (+26 to +31‰) (δ^i O in ‰ = 152 153 $[({}^{i}O/{}^{16}O)_{sample}/({}^{i}O/{}^{16}O)_{VSMOW} - 1] \times 1000; i = 17 \text{ or } 18 \text{ and } VSMOW \text{ is the terrestrial standard material},$ 154 Vienna standard mean ocean water) (Fig. 2a). The simple average of the Δ^{17} O values, the deviation from the terrestrial fractionation line defined as $\Delta^{17}O = \delta^{17}O - 0.52 \times \delta^{18}O$, of the C0002 and Ivuna 155 calcite is $\pm 1.37 \pm 0.40\%$ (2SE, N = 17). The Δ^{17} O values of the C0002, A0058, and Ivuna dolomite 156 157 are systematically lower than those of the calcite, and the average Δ^{17} O value of the dolomite (+0.26 $\pm 0.23\%$, 2SE, N = 16) is closer to the whole-rock values of three Ryugu samples⁸ ($\pm 0.61 \pm 0.28\%$, 158 159 2SD; Fig. 2b). The previously measured O isotope compositions of the A0058 and Ivuna dolomite are 160 in good agreement with those of the C0002 dolomite measured in this study⁸.

161

162 Like the δ^{18} O values, the dolomite in A0058, C0002, and Ivuna has a relatively narrow range of δ^{13} C 163 values from +67 to +75‰ (δ^{13} C in ‰ = [(13 C/ 12 C)_{sample}/(13 C/ 12 C)_{VPDB} – 1] × 1000; VPDB is the 164 terrestrial standard material, Vienna Pee Dee belemnite) (Fig. 3). The δ^{13} C and δ^{18} O values of the 165 dolomite are broadly consistent with the bulk δ^{13} C and δ^{18} O values of carbonates, including calcite,

- dolomite, and breunnerite, in CI chondrites (Ivuna and Orgueil) determined on whole-rock samples¹⁸,
 corroborating the observation that dolomite is the major carbonate mineral. On the other hand, the
- 168 δ^{13} C values of the calcite in C0002 and Ivuna are highly heterogeneous from grain to grain, ranging
- 169 from +65 to +108‰, and they are commonly higher than those of the Ryugu and Ivuna dolomite (Fig.
- 170 3). These variations have also been reported for other Ryugu samples²⁰, confirming that our results
- 171 represent the isotopic characteristics of Ryugu carbonates. For calcite grains on which we conducted
- 172 multiple measurements, the δ^{13} C values in each grain are identical within uncertainties of ~5.1‰. Thus,
- 173 the heterogeneity of δ^{13} C values within individual calcite grains is likely no larger than 10‰, which
- 174 is much smaller than the heterogeneity of >40% between grains.
- 175

176 Isotope signatures of the Ryugu and CI carbonates

177 If the carbonates were in O and C isotopic equilibrium with the aqueous fluid when they precipitated, 178 their O and C isotope compositions would have been determined by the mass dependent equilibrium 179 isotopic fractionation between carbonates and water for O, and that between carbonates and dissolved CO3²⁻ (and other dissolved C-bearing chemical species) for C. The magnitude of this equilibrium 180 isotopic fractionation depends on temperature²¹. Thus, the O and C isotope compositions of the 181 carbonates would reflect those of water and CO_3^{2-} as well as their formation temperatures. Assuming 182 183 equilibrium, the temperature of dolomite-magnetite precipitation in the Ryugu A0058 sample analysed 184 previously is estimated to be 37 ± 10 °C (ref.⁸), while the inferred alteration temperatures of CI 185 chondrites range up to 150 °C (ref.²²). The equilibrium O isotopic fractionation between water and 186 calcite leads to the enhancement of δ^{18} O values in calcite relative to water by approximately +38, +28 and +13‰ at 0, 40, and 150 °C, respectively²³. Thus, the δ^{18} O variation of ~22‰ observed in the 187 188 Ryugu calcite (Fig. 2) is potentially explained by formation temperatures that varied from 0 to 150 °C 189 assuming a fixed δ^{18} O value of water.

190

191 However, this argument does not necessarily mean that variable formation temperature is the sole explanation for the observed δ^{18} O variation of the calcite. Indeed, the lack of a simple correlation 192 between δ^{18} O and δ^{13} C values (Fig. 3) implies that variable formation temperatures alone cannot 193 194 explain the observed δ^{18} O and δ^{13} C variations because the δ^{18} O and δ^{13} C values of carbonates should co-vary with their formation temperatures¹⁸. Rather it seems likely that the δ^{18} O value of water and/or 195 the δ^{13} C value of CO₃²⁻ varied spatially and/or temporally. A previous study used clumped isotope 196 thermometry for carbonates in Mighei-type (CM) chondrites and demonstrated that the δ^{18} O values of 197 198 water are variable between samples¹⁶.

199

200 Mass balance calculations^{24,25} and the O isotope composition of the putative, early solar system 201 water^{26,27} suggest that prior to the onset of alteration, water in the CI and other carbonaceous chondrites

- had a significantly higher Δ^{17} O value, and possibly δ^{18} O value, than the anhydrous silicates²⁸. Thus, 202 as alteration progressed, the Δ^{17} O value, and possibly the δ^{18} O value, of the altering fluid would have 203 decreased^{24,25}. The Δ^{17} O values of the carbonates and fluid will be identical at equilibrium, and thus, 204 205 the Δ^{17} O values of the carbonates are a measure of the degree of progress of water-rock interactions. 206 The Δ^{17} O values of the dolomite in the Ryugu and Ivuna samples are systematically lower than those 207 of the calcite, and the Ryugu and Ivuna carbonates show resolvable Δ^{17} O variations beyond uncertainties (Fig. 2b). Therefore, the calcite with systematically higher Δ^{17} O values formed from less 208 209 "evolved" fluids, and crystallized earlier than the dolomite. The fact that calcite is more prevalent in 210 less-altered areas, as shown by primary anhydrous silicates, suggests that Ca was more easily leached 211 during incipient aqueous alteration than Mg, allowing the formation of calcite before dolomite.
- 212

213 Like O isotopes, the C isotope compositions of carbonates are determined not only by their formation temperatures but also by the δ^{13} C values of dissolved CO₃²⁻. The C isotope composition of CO₃²⁻ 214 215 could have varied due to (i) Rayleigh-type isotopic fractionation as a result of the preferential escape of ¹²C-rich gaseous species like CH₄ (ref.¹⁶), (*ii*) the progressive formation of carbonates, i.e., 216 fractional crystallization, (*iii*) the mixing of two or more C reservoirs with distinct δ^{13} C values that 217 218 supplied CO_3^{2-} (ref.¹⁹), and (*iv*) the change in the chemical speciation of the C-bearing gaseous species, such as CO₂, CO, and CH₄, due to varying O and H partial pressures. Rayleigh-type isotopic 219 220 fractionation was not the primary mechanism for producing the observed δ^{13} C variation because it 221 would have resulted in higher δ^{13} C values in the dolomite, which formed from the more evolved fluids 222 than the calcite, whereas the opposite is observed. The influence of fractional crystallization during 223 carbonate formation was also minimal because the rare calcite that formed early presumably when a larger CO_3^{2-} pool was available displays a larger $\delta^{13}C$ variation than the more common dolomite, 224 225 which is opposite to what one would expect in this scenario.

226

A previous study advocated that the observed δ^{13} C variation resulted from the mixing of C reservoirs with distinct δ^{13} C values reflecting the spatially heterogenous distribution of different C reservoirs¹⁹. Possible C reservoirs include C-bearing gaseous species such as CO₂, CO, and CH₄, originally accreted in ices, and organic matter, and the previous study invoked CO₂-bearing ices as a ¹³C-rich reservoir¹⁹. However, the mixing timescales of gaseous species in the Ryugu/CI parent asteroids would have been short unless the permeability was extremely low. Therefore, the spatially heterogeneous distribution

- 233 of C reservoirs would have not persisted for long. Furthermore, it is unclear why the calcite has
- 234 commonly higher δ^{13} C values than the dolomite, and calcite with lower δ^{13} C values is generally absent.
- 235

236 **Temporal change in oxygen fugacity and gaseous species**

237 Instead of spatial heterogeneity, the temporal variation in the δ^{13} C values of the C reservoirs and their

238 chemical speciation likely occurred due to a change in O partial pressure, or more precisely, O fugacity 239 (fO_2), which is O partial pressure corrected for nonideal gas behaviour. Oxygen fugacity varied along 240 with the production of H₂ via the oxidation of Fe in metal and silicates by H₂O and the subsequent 241 escape of H₂ from the system, perhaps by diffusion or by making fractures in the parent body²⁹. Thus, 242 fO_2 was determined by the relative rates of the production and escape of H₂; in this scenario, fO_2 would

- 243 have at first decreased and then increased. In the case of CM chondrites, the amount of Fe^{3+} in their
- 244 matrices increases with increasing alteration³⁰, which is in line with this scenario.
- 245

The Fe and Mn abundances of terrestrial carbonates that reflect the Fe²⁺ and Mn²⁺ activities in fluids have been used to infer the redox conditions under which they precipitated³¹. However, the Fe and Mn abundances of Ryugu and Ivuna carbonates were likely controlled not only by redox conditions but also by the amounts of these cations leached from primary minerals during the progressive aqueous alteration³². Therefore, the zoning of Fe and Mn observed in the dolomite grains may not be a direct proxy for the temporal change in fO_2 . Instead, here we propose that the $\delta^{13}C$ values of Ryugu and Ivuna carbonates represent a record of such temporal fO_2 variation.

253

To see how the δ^{13} C values of carbonates will change with varying fO_2 , we consider a rather simple 254 model, where gaseous CO_2 and CO and carbonates (and dissolved CO_2 , HCO_3^- , and CO_3^{2-}) are in C 255 256 isotopic equilibrium, and the CO_2/CO ratio increases, corresponding to an increase in fO_2 . In cometary 257 ices, CO₂ and CO are the most abundant C-bearing chemical species³³, and the Ryugu/CI parent bodies 258 would have accreted significant amounts of CO₂- and CO-bearing ices if they formed in the distal 259 solar system^{11,28}. At the earliest stage of aqueous alteration, the CO₂/CO ratio may have been 260 characterized by that of the accreted ices, which may be around unity or higher as observed for 261 cometary ices^{33,34}. We assume that the δ^{13} C value of the bulk gas (CO₂ + CO), δ^{13} C_{bulk}, is constant regardless of the CO₂/CO ratio. Then, the $\delta^{13}C_{\text{bulk}}$ is given by mass balance as $\delta^{13}C_{\text{bulk}} = x \, \delta^{13}C_{\text{CO2}} + 1$ 262 $(1 - x) \delta^{13}C_{CO}$, where x is the mole fraction of CO₂ defined by $x = CO_2/(CO_2 + CO)$, $\delta^{13}C_{CO2}$ and 263 $\delta^{13}C_{CO}$ are the $\delta^{13}C$ values of CO₂ and CO, respectively. Thus, using Δ defined by $\delta^{13}C_{CO2} - \delta^{13}C_{CO}$, 264 $\delta^{13}C_{CO2}$ and $\delta^{13}C_{CO}$ are given by $\delta^{13}C_{CO2} = \delta^{13}C_{bulk} + (1-x)\Delta$ and $\delta^{13}C_{CO} = \delta^{13}C_{bulk} - x\Delta$, respectively. 265 266 The Δ value is approximated by 1000 ln α , where α is the C isotopic fractionation factor between CO₂ and CO defined by $\binom{13}{2} \binom{12}{3} \binom{13}{2} \binom$ 267 at 0, 40, and 150 °C, respectively^{21,35}). Therefore, with increasing x (and fO_2), both the δ^{13} C values of 268 269 CO_2 and CO will decrease monotonically, and the $\delta^{13}C$ values of carbonates will also decrease. Thus, 270 in this simple model, a δ^{13} C variation of carbonates comparable to the observation (~40‰) is expected 271 if the x value varies from 0.5 to 1 (Fig. 3). The redox states evolved within a few million years after the birth of the solar system as inferred from the ⁵³Mn-⁵³Cr chronometry of Ryugu/CI dolomite^{8,10,20,36}. 272 273 The presence of CH₄, another possible reducing gas, does not change this conclusion because the

behaviour of CH_4 in terms of C isotopic fractionation against CO_2 is quite similar to that of CO (ref.²¹). Our model where CO_2 and CO are in C isotopic equilibrium at low temperatures requires the presence of as yet unidentified processes/catalysts. The isotopic equilibrium in thermally matured natural gases has been recently discussed³⁷.

278

279 The characteristic δ^{13} C values of Ryugu and Ivuna carbonates are not observed for other aqueously altered meteorites such as CM chondrites and the ungrouped carbonaceous chondrite Tagish 280 Lake^{19,38,39}, suggesting that the redox conditions as well as the thermal history and accreted materials 281 are unique to their parent bodies. For instance, the δ^{13} C values of CM calcite are variable like Ryugu/CI 282 283 calcite, but the highest reported value (~+80‰) in CM calcite is lower than that of Ryugu/CI calcite 284 (+108‰) (Extended Data Fig. 2). Furthermore, the δ^{13} C values of CM dolomite are also variable from ~+40 to ~+60‰, which is in contrast to the homogeneous δ^{13} C values observed for Ryugu/CI dolomite. 285 Iron in CI chondrites is dominated by octahedral Fe³⁺ and indicates more oxidized conditions than CM 286 chondrites³⁰, which is consistent with the homogeneous δ^{13} C values of Ryugu/CI dolomite. 287

288

289 Because the δ^{13} C values of the Ryugu/CI dolomite are commonly lower and more homogeneous than 290 those of the calcite, the dolomite likely formed at higher fO₂ and/or temperature than the calcite. This, 291 combined with the O isotope signatures of the carbonates, implies that the calcite formed during 292 prograde alteration over wide ranges of fO_2 and temperature, whereas the dolomite formed later during 293 retrograde cooling when the aqueous fluids and silicates approached O isotopic equilibrium. Therefore, 294 when the dolomite formed, the fO_2 was likely high enough that the major gaseous C reservoir was CO_2 295 (i.e., $x \sim 1$). If correct, the δ^{13} C value of CO₂ in C isotopic equilibrium with the dolomite, which has 296 the average δ^{13} C value of ~+70‰, would equal the δ^{13} C value of the bulk gas. Using the C isotopic 297 fractionation factor between dolomite and CO₂, (¹³C/¹²C)_{dolomite}/(¹³C/¹²C)_{CO2}, of ~1.0092 at 40 °C or 298 ~0.9951 at 150 °C (refs.^{40,41}), the $\delta^{13}C_{bulk}$ (= $\delta^{13}C_{CO2}$) value is estimated to be between +61‰ and 299 +75‰.

300

The above simple model implies a δ^{13} C value higher than +60‰ for C-bearing molecules originally 301 302 accreted in ices. Such ¹³C-rich compositions are not common among solar system materials other than meteoritic carbonates and water-soluble organic compounds⁴². A similar level of ¹³C-enrichment ($\delta^{13}C$ 303 = $+65 \pm 51\%$), albeit with large uncertainty, has been reported for CO₂ in the coma of 67P/Churyumov-304 305 Gerasimenko⁴³. The mechanism to produce ¹³C-rich compositions is not well understood, but self-306 shielding during CO photodissociation in the solar nebula or the parent molecular cloud of the solar system is a possible mechanism⁴⁴⁻⁴⁶. Thus, the inferred ¹³C-rich composition of C-bearing molecular 307 ices would have resulted from such physicochemical reactions^{47,48}, and we concluded that the 308 309 Ryugu/CI parent bodies accreted materials that originated from these cold environments.

312

311 Acknowledgements

313 materials for SIMS measurements, and Akira Tsuchiyama for discussion. Hayabusa2 was developed 314 and built under the leadership of Japan Aerospace Exploration Agency (JAXA), with contributions 315 from the German Aerospace Center (DLR) and the Centre National d'Études Spatiales (CNES), and 316 in collaboration with NASA, and other universities, institutes, and companies in Japan. The curation 317 system was developed by JAXA in collaboration with companies in Japan. This research was 318 supported in part by the JSPS KAKENHI grant numbers 19H00725 and 20K20934 to W.F. and T.N. 319 320 **Author contributions** 321 W.F., N.K., K.N., N.S, and H.Y. designed this study and analysed the samples. W.F., N.K., K.N., C.A., 322 and H.Y. were involved in data reduction and data interpretation. W.F. wrote the paper with support 323 and approval of all co-authors. 324 325 **Competing interests** 326 The authors declare no competing interests. 327

We thank Drae Rogers, Michael Spicuzza, and John Valley for the preparation of carbonate standard

Samples	δ ¹⁸ O (‰)	2σ	δ ¹⁷ O (‰)	2σ	$\Delta^{17}O$ (‰)	2σ	δ ¹³ C (‰)	2σ	
Ryugu C0002, calcite									
<i>Calcite</i> #1 <i>a</i>	23.5	0.8	12.5	1.6	0.2	1.5	98.1	4.9	
<i>Calcite</i> #1 <i>b</i>	23.5	0.8	12.5	1.6	0.2	1.5	92.7	4.8	
Calcite#1ave.	23.5	0.8	12.5	1.6	0.2	1.5	95.4	3.4	
Calcite#2	38.4	0.6	22.2	1.1	2.2	1.1	101.7	4.8	
Calcite#3-1	44.3	0.8	24.6	1.4	1.6	1.5	89.9	5.2	
Calcite#3-2a	46.3	0.8	25.9	1.6	1.9	1.7	76.2	5.7	
Calcite#3-2b	46.3	0.8	25.9	1.6	1.9	1.7	84.5	6.5	
Calcite#3-2c	46.3	0.8	25.9	1.6	1.9	1.7	81.0	5.9	
Calcite#3-2ave.	46.3	0.8	25.9	1.6	1.9	1.7	80.6	3.5	
Calcite#3-3	44.5	1.2	24.9	1.5	1.8	1.4	84.1	6.2	
Calcite#3-4	42.9	1.2	23.3	1.3	1.0	1.4	87.0	5.6	
Calcite#4-1	n.d.		n.d.		n.d.		88.2	5.2	
Calcite#4-2	n.d.		n.d.		n.d.		86.1	4.8	
Calcite#5a	41.1	1.2	24.2	1.4	2.9	1.4	93.9	4.8	
Calcite#5b	40.9	1.2	23.2	1.3	1.9	1.4	93.9	4.8	
Calcite#5ave.	41.0	0.8	23.7	1.0	2.4	1.0	93.9	4.8	
			Ivuna, ca	lcite					
Calcite#1	41.6	1.0	22.5	1.3	0.9	1.4	101.5	4.1	
Calcite#2	41.5	1.0	23.4	1.3	1.8	1.4	104.8	4.8	
Calcite#3	40.7	1.1	21.9	1.6	0.8	1.4	107.1	4.7	
Calcite#5	33.9	1.0	20.0	1.3	2.4	1.4	81.4	3.8	
Calcite#6	41.1	1.0	24.0	1.4	2.6	1.4	103.9	6.0	
Calcite#7	38.0	1.0	20.2	1.4	0.4	1.4	98.0	5.1	
Calcite#8-1	37.6	1.0	20.6	1.3	1.1	1.4	79.5	4.4	
Calcite#8-2	37.0	1.0	20.0	1.3	0.8	1.4	89.0	5.2	
Calcite#9a	36.0	1.0	20.2	1.3	1.5	1.4	106.6	5.8	
Calcite#9b	36.0	1.0	20.2	1.3	1.5	1.4	109.9	5.1	
<i>Calcite</i> #9c	36.0	1.0	20.2	1.3	1.5	1.4	106.7	4.8	
Calcite#9ave.	36.0	1.0	20.2	1.3	1.5	1.4	107.8	3.0	
Calcite#10a	43.6	0.6	22.3	1.5	-0.4	1.4	65.4	4.7	
Calcite#10b	40.9	0.7	21.2	1.2	-0.1	1.2	65.4	4.7	
Calcite#10ave.	42.2	0.5	21.7	1.0	-0.3	0.9	65.4	4.7	

Table 1. δ^{18} O, δ^{17} O, Δ^{17} O, and δ^{13} C values of carbonates in Ryugu and Ivuna samples.

-		R	yugu A005	8, dolomi	te					
Dolomite#5	(27.0)	(0.9)	(14.2)	(1.0)	(0.2)	(0.9)	67.7	1.5		
Dolomite#5	(27.0)	(0.9)	(14.2)	(1.0)	(0.2)	(0.9)	67.2	1.2		
Dolomite#5ave.	(27.0)	(0.9)	(14.2)	(1.0)	(0.2)	(0.9)	67.5	1.0		
Dolomite#2	(29.0)	(1.0)	(15.0)	(1.0)	(0.0)	(0.9)	74.8	1.5		
Dolomite#4	(30.0)	(0.9)	(15.8)	(1.0)	(0.2)	(0.9)	72.1	1.3		
Dolomite#1	(29.9)	(0.9)	(14.8)	(1.0)	(-0.7)	(0.9)	71.3	1.2		
Dolomite#3	(29.5)	(0.8)	(15.7)	(1.0)	(0.4)	(0.9)	74.5	1.5		
Ryugu C0002, dolomite										
Dolomite#3a	27.3	1.0	15.2	1.3	1.0	0.9	72.1	1.3		
Dolomite#3b	28.1	1.0	14.7	1.3	0.1	0.9	72.1	1.3		
Dolomite#3ave.	27.7	0.7	14.9	0.9	0.6	0.6	72.1	1.3		
Dolomite#4	28.3	1.0	15.3	1.3	0.6	0.9	70.1	1.3		
Dolomite#1a	28.6	1.0	15.6	1.3	0.7	0.9	68.1	1.3		
Dolomite#1b	28.7	1.0	15.1	1.3	0.2	0.9	68.1	1.3		
Dolomite#1ave.	28.7	0.7	15.4	0.9	0.5	0.6	68.1	1.3		
Dolomite#2a	26.7	1.0	14.1	1.3	0.2	0.9	69.4	1.3		
Dolomite#2b	27.7	1.0	14.1	1.3	-0.3	0.9	69.5	1.3		
Dolomite#2ave.	27.2	0.7	14.1	0.9	0.0	0.6	69.4	0.9		
			Ivuna, de	olomite						
Dolomite#1	(29.0)	(0.7)	(15.6)	(0.8)	(0.5)	(0.9)	68.1	1.3		
Dolomite#7	(30.8)	(0.7)	(16.0)	(0.8)	(0.0)	(0.7)	68.6	1.1		
Dolomite#6	(29.1)	(0.8)	(15.3)	(0.8)	(0.2)	(0.9)	72.3	1.2		
Dolomite#5	(28.8)	(0.8)	(16.3)	(0.8)	(1.3)	(0.9)	67.2	1.2		
Dolomite#4	(26.5)	(0.9)	(14.4)	(0.9)	(0.7)	(1.0)	68.3	1.3		
Dolomite#2a	(29.6)	(0.9)	(15.1)	(0.9)	(-0.3)	(0.8)	67.4	1.3		
Dolomite#2b	(29.6)	(0.9)	(15.1)	(0.9)	(-0.3)	(0.8)	65.6	1.3		
Dolomite#2c	(29.6)	(0.9)	(15.1)	(0.9)	(-0.3)	(0.8)	67.4	1.3		
Dolomite#2ave.	(29.6)	(0.9)	(15.1)	(0.9)	(-0.3)	(0.8)	66.8	0.7		
Dolomite#3	(29.2)	(1.0)	(15.4)	(1.0((0.2)	(1.1)	68.8	1.2		

n.d.: Not determined.

The O isotope compositions of the Ryugu A0058 and Ivuna dolomite, shown in parentheses, are taken from Yokoyama et al. (2022) (ref.⁸)

 2σ errors are either external reproducibility (2SD) of standard measurements or internal precision (2SE) of the data within single measurements, whichever is the largest.

In case that we analysed single grains multiple times, as shown in Italic, we averaged the data and calculated the corresponding uncertainties by propagating individual errors.





Fig. 1. Calcite grains in the Ryugu C0002 sample. (a) Mg (red), Si (green), and Ca (blue) elemental maps showing the distribution of calcite grains. Calcite grains (blue) in the Ryugu and Ivuna samples can be found in limited areas with Mg-rich olivine (yellow) and pyroxene (light green). Purple grains are dolomite. The large red grain is breunnerite. (b) Backscattered electron image (BEI) of a calcite grain.





337 Fig. 2. Oxygen isotope compositions of the calcite and dolomite in Ryugu and Ivuna samples. The 338 whole-rock O isotope compositions of the Ryugu A0107 and C0002 samples are also shown for 339 reference⁸. The changes in O isotope compositions due to variable formation temperatures and waterrock interaction are illustrated by arrows. (a) Oxygen three-isotope plot showing δ^{18} O vs. δ^{17} O values. 340 TFL: Terrestrial fractionation line defined as $\delta^{17}O = 0.52 \times \delta^{18}O$. (b) $\delta^{18}O$ vs. $\Delta^{17}O$ values. Data are 341 presented as mean values $\pm 2\sigma$ errors which are either external reproducibility (2SD, N = 6-20342 343 depending on the measurement sessions) of standard measurements or internal precision (2SE) of the 344 data within single measurements, whichever is the largest.





346 Fig. 3. Comparison between C and O isotope compositions of the calcite and dolomite in Ryugu 347 and Ivuna samples. Errors are 2σ (see also Table 1). The values of carbonates measured for wholerock CI chondrites (Ivuna and Orgueil) are also shown for reference¹⁸. The changes in C and O isotope 348 349 compositions due to variable formation temperatures and O fugacity, and water-rock interaction are 350 illustrated by arrows. The calcite shows much larger variations in both C and O isotope compositions than the dolomite. Note that no simple correlation between $\delta^{13}C$ and $\delta^{18}O$ values can be seen. Data are 351 presented as mean values $\pm 2\sigma$ errors which are either external reproducibility (2SD, N = 6-20352 353 depending on the measurement sessions) of standard measurements or internal precision (2SE) of the 354 data within single measurements, whichever is the largest. 355

356 **References:**

- Tachibana, S. et al. Hayabusa2: Scientific importance of samples returned from C-type near-Earth
 asteroid (162173) 1999 JU₃. *Geochem. J.* 48, 571-587 (2014).
- Tachibana, S. et al. Pebbles and sand on asteroid (162173) Ryugu: In situ observation and
 particles returned to Earth. *Science* 375, 1011-1016 (2022).
- Binzel, R. P., Harris, A. W., Bus, S. J. & Burbine, T. H. Spectral properties of near-Earth objects:
 Palomar and IRTF results for 48 objects including spacecraft targets (9969) Braille and (10302)
 1989 ML. *Icarus* 151, 139-149 (2001).
- Campins, H. et al. Spitzer observations of spacecraft target 162173 (1999 JU3). *Astron. Astrophys.* 503, L17–L20 (2009).
- 366 5. Watanabe, S. et al. Hayabusa2 arrives at the carbonaceous asteroid 162173 Ryugu—A spinning
 367 top-shaped rubble pile. *Science* 364, 268-272 (2019).
- 368 6. Sugita, S. et al. The geomorphology, color, and thermal properties of Ryugu: Implications for
 369 parent-body processes. *Science* 364, eaaw0422 (2019).
- 370 7. Kitazato K. et al. The surface composition of asteroid 162173 Ryugu from Hayabusa2 near371 infrared spectroscopy. *Science* 364, 272-275 (2019).
- 372 8. Yokoyama, T. et al. The first returned samples from a C-type asteroid show kinship to the
 373 chemically most primitive meteorites. *Science* 379, eabn7850 (2022).
- 9. Nakamura, T. et al. Formation and evolution of carbonaceous asteroid Ryugu: Direct evidence
 from returned samples. *Science*, 10.1126/science.abn8671 (2022).
- 10. Nakamura, E. et al. On the origin and evolution of the asteroid Ryugu: A comprehensive
 geochemical perspective. *Proc. Jpn. Acad., Ser. B* 98, 227-282 (2022).
- 11. Hopp, T. et al. Ryugu's nucleosynthetic heritage from the outskirts of the Solar System. *Sci. Adv.*8, eadd8141 (2022).
- Paquet, M. et al. Contribution of Ryugu-like material to Earth's volatile inventory by Cu and Zn
 isotopic analysis. *Nat. Astron.* 7, 182-189 (2023).
- 382 13. Moynier, F. et al. The Solar System calcium isotopic composition inferred from Ryugu samples.
 383 *Geochem. Persp. Let.* 24, 1-6 (2022).
- Johnson, C. A. & Prinz, M. Carbonate compositions in CM and CI chondrites, and implications
 for aqueous alteration. *Geochim. Acta* 57, 2843-2852 (1993).
- 15. Riciputi, L. R., McSween, H. Y. Jr., Johnson, C. A. & Prinz, M. Minor and trace element
 concentrations in carbonates of carbonaceous chondrites, and implications for the compositions
 of coexisting fluids. *Geochim. Cosmochim. Acta* 58, 1343-1351 (1994).
- 389 16. Guo, W. & Eiler, J. M. Temperatures of aqueous alteration and evidence for methane generation
 390 on the parent bodies of the CM chondrites. *Geochim. Cosmochim. Acta* **71**, 5565-5575 (2007).
- 391 17. Verdier-Paoletti, M. J. et al. Oxygen isotope constraints on the alteration temperatures of CM

- 392 chondrites. *Earth Planet. Sci. Lett.* 458, 273-281 (2017).
- 18. Alexander, C. M. O'D., Bowden, R., Fogel, M. L. & Howard, K. T. Carbonate abundances and
 isotopic compositions in chondrites. *Meteorit. Planet. Sci.* 50, 810-833 (2015).
- Fujiya, W. et al. Migration of D-type asteroids from the outer solar system inferred from
 carbonate in meteorites. *Nat. Astron.* 3, 910-915 (2019).
- McCain, K. A. et al. Early fluid activity on Ryugu inferred by isotopic analyses of carbonates and
 magnetite. *Nat. Astron.***7**, 309-317 (2023).
- 21. Chacko, T., Cole, D. R. & Horita, J. Equilibrium oxygen, hydrogen and carbon isotope
 fractionation factors applicable to geologic systems. In *Stable Isotope Geochemistry* (eds. Valley,
 J. W. and Cole, D. R.) 1-81 (Mineralogical Society of America, 2001).
- 22. Zolensky, M. E., Bourcier, W. L. & Gooding, J. L. Aqueous alteration on the hydrous asteroids:
 Results of EQ3/6 computer simulations. *Icarus* 78, 411-425 (1989).
- 23. Zheng, Y.-F. On the theoretical calculations of oxygen isotope fractionation factors for carbonatewater systems. *Geochem. J.* 45, 341-354 (2011).
- 24. Clayton, R. N. & Mayeda, T. K. The oxygen isotope record in Murchison and other carbonaceous
 chondrites. *Earth Planet. Sci. Lett.* 67, 151-161 (1984).
- 408 25. Marrocchi, Y., Bekaert, D. V. & Piani, L. Origin and abundance of water in carbonaceous
 409 asteroids. *Earth Planet. Sci. Lett.* 482, 23-32 (2018).
- 26. Sakamoto, N. et al. Remnants of the early Solar System water enriched in heavy oxygen isotopes. *Science* 317, 231-233 (2007).
- 412 27. Vacher, L. G., Marrocchi, Y., Verdier-Paoletti, M. J., Villeneuve, J. & Gounelle, M. Inward radial
 413 mixing of interstellar water ices in the solar protoplanetary disk. *Astrophys. J.* 827, L1 (2016).
- 414 28. Kawasaki, N. et al. Oxygen isotopes of anhydrous primary minerals show kinship between steroid
 415 Ryugu and comet 81P/Wild2. *Sci. Adv.* 8, eade2067 (2022).
- Wilson L., Keil, K., Browning, L. B., Krot, A. N. & Bourcier, W. Early aqueous alteration,
 explosive disruption, and reprocessing of asteroids. *Meteorit. Planet. Sci.* 34, 541-557 (1999).
- 30. Beck, P. et al. The redox state of iron in the matrix of CI, CM and metamorphosed CM chondrites
 by XANES spectroscopy. *Geochim. Cosmochim. Acta* 99, 305-316 (2012).
- 31. Barnaby, R. J. & Rimstidt, J. D. Redox conditions of calcite cementation interpreted from Mn
 and Fe contents of authigenic calcites. *Geol. Soc. Am. Bull.* 101, 795-804.
- 422 32. Fujiya, W., Aoki, Y., Ushikubo, T., Hashizume, K. & Yamaguchi, A. Carbon isotopic evolution
 423 of aqueous fluids in CM chondrites: Clues from *in-situ* isotope analyses within calcite grains in
 424 Yamato-791198. *Geochim. Cosmochim. Acta* 274, 246-260 (2020).
- 33. Mumma, M. J. & Charnley, S. B. The chemical composition of comets—Emerging taxonomies
 and natal heritage. *Annu. Rev. Astron. Astrophys.* 49, 471–524 (2011).
- 427 34. Ootsubo, T. et al. AKARI Near-infrared spectroscopic survey for CO₂ in 18 comets. *Astrophys.*

428 *J.* **752**, 15 (2012).

- 35. Richet, P., Bottinga, Y. & Javoy, M. A review of hydrogen, carbon, nitrogen, and chlorine stable
 isotope fractionation among gaseous molecules. *Ann. Rev. Earth Planet. Sci.* 5, 65-110 (1977).
- 431 36. Fujiya, W., Sugiura, N., Sano, Y. and Hiyagon, H. Mn–Cr ages of dolomites in CI chondrites and 432 the Tagish Lake ungrouped carbonaceous chondrite. *Earth Planet. Sci. Lett.* **362**, 130-142 (2013).
- 433 37. Thiagarajan, N. et al. Isotopic evidence for quasi-equilibrium chemistry in thermally mature
 434 natural gases. *Proc. Natl Acad. Sci. USA* 117, 3989-3995 (2017).
- 38. Telus, M., Alexander, C. M. O'D., Hauri, E. H. & Wang, J. Calcite and dolomite formation in the
 CM parent body: Insight from *in-situ* C and O isotope analyses. *Geochim. Cosmochim. Acta* 260,
 275-291 (2019).
- 438 39. Vacher, L. G., Marrocchi, Y., Villeneuve, J., Verdier-Paoletti, M. J. & Gounelle, M. Petrographic
 439 and C & O isotopic characteristics of the earliest stages of aqueous alteration of CM chondrites.
 440 *Geochim. Cosmochim. Acta* 213, 271-290 (2017).
- 441 40. Sheppard, S. M. F. & Schwarcz, H. P. Fractionation of carbon and oxygen isotopes and
 442 magnesium between coexisting metamorphic calcite and dolomite. *Contr. Mineral. Petrol.* 26,
 443 161-198 (1970).
- 444 41. Romanek, C. S., Grossman, E. L. & Morse, J. W. Carbon isotopic fractionation in synthetic
 445 aragonite and calcite: Effects of temperature and precipitation rate. *Geochim. Cosmochim. Acta*446 56, 419-430 (1992).
- 42. Aponte, J. C., McLain, H. L., Dworkin, J. P. & Elsila, J. E. Aliphatic amines in Antarctic CR2,
 CM2, and CM1/2 carbonaceous chondrites. *Geochim. Cosmochim. Acta* 189, 296-311 (2016).
- 43. Hässig, M. et al. Isotopic composition of CO₂ in the coma of 67P/Churyumov-Gerasimenko
 measured with ROSINA/DFMS. *Astron. Astrophys.* 605, A50 (2017).
- 44. Yurimoto, H. & Kuramoto, K. Molecular cloud origin for the oxygen isotope heterogeneity in the
 solar system. *Science* **305**, 1763-1766.
- 45. Lyons, J. R. & Young, E. D. CO self-shielding as the origin of oxygen isotope anomalies in the
 early solar nebula. *Nature* 435, 317-320.
- 455 46. Lyons, J. R., Gharib-Nezhad, E. & Ayres, T. R. A light carbon isotope composition for the Sun.
 456 *Nat. Commun.* 9, 908 (2018).
- 457 47. Visser, R., van Dishoeck, E. F. & Black, J. H. The photodissociation and chemistry of CO
 458 isotopologues: applications to interstellar clouds and circumstellar disks. *Astron. Astrophys.* 503,
 459 323-353 (2009).
- 460 48. Woods, P. M. & Willacy, K. Carbon isotope fractionation in protoplanetary disks. *Astrophys. J.*461 693, 1360-1378 (2009).

463 Methods

464 **Isotope measurement using ion microprobe**

465 We produced the polished sections of Ryugu samples A0058-C1001 and C0002-C1001, and Ivuna embedded in $epoxy^{8,28}$. The polished sections were coated with a thin (~5 nm) gold film using a Leica 466 467 EM ACE600 coater at Hokkaido University for backscattered electron (BSE) and X-ray imaging, and 468 elemental analysis before *in-situ* O and C isotope measurements. We observed their mineralogy and 469 petrology and located carbonate grains using a field-emission scanning electron microscope (FE-SEM; 470 JEOL JSM-7000F) equipped with an energy dispersive X-ray spectrometer (EDS; Oxford X-Max 150) 471 at Hokkaido University. The beam currents were ~ 2 nA and ~ 1 nA for the X-ray mapping and 472 quantitative analysis, respectively. Quantitative calculations were conducted using Oxford AZtec 473 software.

474

We selected five and four dolomite grains from the Ryugu samples A0058 and C0002, respectively, and nine calcite grains from C0002 for isotope analyses. We also analysed seven dolomite grains and ten calcite grains from Ivuna. Before the isotope analyses, the samples were coated again with an additional thin (~65 nm) gold film.

479

480 We conducted the O and C isotope measurements of the selected carbonate grains using secondary ion 481 mass spectrometry (SIMS; CAMECA ims-1280HR) at Hokkaido University. Instrumental mass fractionation (IMF) was corrected using the UWC3 calcite standard and a series of dolomite-ankerite 482 483 standards from the WiscSIMS laboratory for calcite and dolomite, respectively⁴⁹⁻⁵¹. Measurement 484 spots were observed using the FE-SEM after the SIMS measurements and data from spots with 485 inclusions or overlapping matrix minerals were rejected. The reported uncertainties (2σ) of isotope 486 compositions are the larger of the external reproducibility, i.e., 2 standard deviation (2SD) of standard 487 measurements or internal precision, i.e., 2 standard error (2SE) of data within single measurements of 488 unknown samples.

489

The procedures of O isotope measurement of dolomite were described by a previous study⁸. For O 490 491 isotope measurement of calcite, as previously described for the O isotope measurement of olivine²⁸, secondary ${}^{16}O^{-}$, ${}^{17}O^{-}$, and ${}^{18}O^{-}$ ions produced by a Cs⁺ primary ion beam (~30 pA, ~3 µm) were 492 simultaneously collected using Faraday cup (FC: $10^{11} \Omega$), electron multiplier (EM), and EM detectors, 493 494 respectively. Mass resolving power was >6000 sufficient to resolve ${}^{17}O^{-}$ from ${}^{16}OH^{-}$. The secondary ion intensities of ${}^{16}\text{O}^-$ were 2-3 \times 10⁷ cps. The measurement time was 240 seconds. The ${}^{16}\text{OH}^-$ count 495 rate was measured immediately after each measurement, and we made a small tail correction on ¹⁷O⁻; 496 497 its contribution to ${}^{17}\text{O}^-$ was typically less than ~0.1‰ and up to ~0.5‰ for a few analyses. The typical 498 uncertainties of δ^{17} O, δ^{18} O, and Δ^{17} O values were 1.4, 0.9, and 1.4‰, respectively.

500 For C isotope measurement of dolomite, secondary ${}^{12}C^{-}$ and ${}^{13}C^{-}$ ions produced by a Cs⁺ primary ion 501 (~50 pA, ~2 µm) were simultaneously collected using FC (10¹² Ω) and EM detectors, respectively. We 502 scanned the primary ion beam across 1 × 1 µm-sized areas to make the SIMS pits shallower and 503 suppress ratio drifts during measurements. Mass resolving power was ~4500, sufficient to resolve ${}^{13}C^{-}$ 504 from ${}^{12}CH^{-}$. The secondary ion intensities of ${}^{12}C^{-}$ were 5-6 × 10⁵ cps. The measurement time was 480

- 505 seconds. The typical uncertainty of δ^{13} C values was 1.3‰.
- 506

507 For C isotope measurement of calcite, secondary ${}^{12}C^{-}$ and ${}^{13}C^{-}$ ions produced by a Cs⁺ primary ion 508 (~3 pA, ~1 µm) were simultaneously collected using two EM detectors. We scanned the primary ion 509 beam on 1 × 1 µm-sized areas. Mass resolving power was ~4500. The secondary ion intensities of ${}^{12}C^{-}$ 510 were ~2 × 10⁴ cps. The measurement time was 800 seconds. The typical uncertainty of $\delta^{13}C$ values 511 was 5.1‰. The larger uncertainty of the calcite measurement was due to the smaller primary ion beam 512 intensity than for the dolomite measurement to analyse small calcite grains.

513

514 Data availability

515 All data generated or analysed during this study are included in this published article (and its 516 supplementary information files) and are available via Zenodo 517 (https://doi.org/10.5281/zenodo.7957625). As the initial analysis of Ryugu samples collected by the 518 Hayabusa2 spacecraft, the specimens analysed in this study were allocated to us by JAXA. The Ivuna 519 specimen used in this study was kindly proved by the Natural History Museum, UK.

520

521 **References:**

- 49. Kozdon, R., Ushikubo, T., Kita, N. T., Spicuzza, M. & Valley, J. W. Intratest oxygen isotope
 variability in the planktonic foraminifer *N. pachyderma*: Real vs. apparent vital effects by ion
 microprobe. *Chem. Geol.* 258, 327-337 (2009).
- 525 50. Śliwiński, M. G. et al. Secondary ion mass spectrometry bias on isotope ratios in dolomite– 526 ankerite, Part I: δ^{18} O matrix effects. *Geostand. Geoanal. Res.* **40**, 157-172 (2015a).

527 51. Śliwiński, M. G. et al. Secondary ion mass spectrometry bias on isotope ratios in dolomite-

- 528 ankerite, Part II: δ^{13} C matrix effects. *Geostand. Geoanal. Res.* **40**, 173-184 (2015b).
- 529
- 530