



The University of Manchester Research

Insights into the origin of carbonaceous chondrite organics from their triple oxygen isotope composition

DOI: 10.1073/pnas.1808101115

Document Version

Accepted author manuscript

Link to publication record in Manchester Research Explorer

Citation for published version (APA):

Tartese, R., Chaussidon, M., Gurenko, A., Delarue, F., & Robert, F. (2018). Insights into the origin of carbonaceous chondrite organics from their triple oxygen isotope composition. *Proceedings of the National Academy of Sciences of the United States of America*. https://doi.org/10.1073/pnas.1808101115

Published in:

Proceedings of the National Academy of Sciences of the United States of America

Citing this paper

Please note that where the full-text provided on Manchester Research Explorer is the Author Accepted Manuscript or Proof version this may differ from the final Published version. If citing, it is advised that you check and use the publisher's definitive version.

General rights

Copyright and moral rights for the publications made accessible in the Research Explorer are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

Takedown policy

If you believe that this document breaches copyright please refer to the University of Manchester's Takedown Procedures [http://man.ac.uk/04Y6Bo] or contact uml.scholarlycommunications@manchester.ac.uk providing relevant details, so we can investigate your claim.



1	Insights into the origin of carbonaceous chondrite organics from their triple
2	oxygen isotope composition
3	
4	Romain Tartèse ^{a,*} , Marc Chaussidon ^b , Andrey Gurenko ^c , Frédéric Delarue ^d , François Robert ^e
5	
6	
7	^a School of Earth and Environmental Sciences, The University of Manchester, Manchester, M13
8	9PL, UK.
9	^b Institut de Physique du Globe de Paris (IPGP), Université Sorbonne-Paris-Cité, Université Paris
10	Diderot, CNRS UMR 7154, Paris, France
11	^c Centre de Recherches Pétrographiques et Géochimiques, UMR 7358, Université de Lorraine,
12	54501 Vandoeuvre-lès-Nancy, France.
13	^d Sorbonne Université, UPMC, CNRS, EPHE, PSL, UMR 7619 METIS, 4 place Jussieu, F-75005
14	Paris, France.
15	^e Institut de Minéralogie, de Physique des Matériaux et de Cosmochimie, Muséum National
16	d'Histoire Naturelle, Sorbonne Universités, CNRS, UPMC & IRD, 75005 Paris, France
17	
18	*corresponding author: romain.tartese@manchester.ac.uk – Tel: +44161 275 6908
19	
20	Short title: O isotope composition of carbonaceous chondrite organics.
21	To be submitted to PNAS
22	Categories: Physical Sciences > Earth, Atmospheric, and Planetary Sciences
23	

24 Abstract

Dust grains of organic matter were the main reservoir of C and N in the forming Solar System, 25 26 and are thus considered as an essential ingredient for the emergence of life. Yet, the physical environment and the chemical mechanisms at the origin of these organic grains are still highly 27 debated. In this study we report the first high precision triple oxygen isotope composition for 28 insoluble organic matter isolated from three emblematic carbonaceous chondrites, Orgueil, 29 30 Murchison, and Cold Bokkeveld. These results suggest that the O isotope composition of carbonaceous chondrite insoluble organic matter falls on a slope 1 correlation line in the triple 31 32 oxygen isotope diagram. The lack of detectable mass-dependent O isotopic fractionation, indicated by the slope 1 line, suggest that the bulk of carbonaceous chondrite organics did not 33 34 form on asteroidal parent bodies during low temperature hydrothermal events. On the other hand, these new O isotope data, together with the H and N isotope characteristics of insoluble organic 35 36 matter, may indicate that parent bodies of different carbonaceous chondrite types largely accreted 37 organics formed locally in the protosolar nebula, possibly by photochemical dissociation of C-38 rich precursors.

39

40 Keywords

41 Carbonaceous chondrites; Organic matter; Oxygen isotopes; Protosolar nebula; Secondary ion
42 mass spectrometry.

43

44 Significance Statement

Refractory organic matter found in volatile-rich asteroidal materials essentially comprise the 45 elements C, H, O, N and S, which are thought to be important building blocks for life. 46 Characterizing the origin(s) of these organics thus constitutes a key step to constrain the origin of 47 life on Earth and appraise the habitability potential of other worlds. Yet how and where these 48 organics formed is still highly debated. In this study we have determined the oxygen isotope 49 composition of refractory organics from two families of carbonaceous chondrites. These data 50 51 suggest that these organics formed in the nascent Solar System, possibly through chemical reactions occurring in the disk surrounding the young Sun. 52

53

54 \body

55

56 Introduction

Type 1-2 carbonaceous chondrites (CC) contain several wt.% carbon that mostly occurs as small 57 patches of organic matter (OM) dispersed in the fine-grained matrix (1). Because this OM 58 59 possibly played a key role in the development of life on the early Earth, its molecular structure, 60 and its chemical and isotopic compositions have been extensively investigated (see 2 and 61 references therein for a recent review). Despite this profusion of structural, chemical, and isotopic 62 information, the question of whether CC OM formed in the cold interstellar medium (e.g., 3-5), in the protosolar nebula (PSN) (e.g., 6-9), or is a product of organic synthesis during 63 hydrothermalism on CC parent bodies (e.g., 10) remains highly debated, notably because the 64 65 extent of chemical and isotopic alteration of OM during secondary processes on CC parent bodies is debated (11-15). 66

67

Oxygen is the third most abundant element in the Solar System and has three stable isotopes ¹⁶O, ¹⁷O and ¹⁸O. Because different fractionation laws govern interplanetary and intraplanetary processes (e.g., 16), the oxygen three-isotope system can provide information that cannot be accessed using other two-isotope systems of light elements such as H, C and N. In planetary bodies, variations of the ¹⁷O/¹⁶O and ¹⁸O/¹⁶O ratios almost always obey the mass-dependent relationship $\delta^{17}O \sim 0.52 \times \delta^{18}O^1$, while oxygen isotope abundance variations between Solar System gas and solids are primarily governed by the mass-independent relationship $\delta^{17}O \sim 1.0 \times$

¹this δ -notation represents deviations in parts per thousand (‰) of the ^{17,18}O/¹⁶O ratios relative to those of the Standard Mean Ocean Water (SMOW), according to the equation $\delta^{17,18}O = [(^{17,18}O/^{16}O)_{sample}/(^{17,18}O/^{16}O)_{SMOW} - 1] \times 1000).$

 δ^{18} O (15). Much of our understanding of how our Solar System formed and evolved is thus based on O isotope studies of meteoritic materials (e.g., 16-17), and this should apply to carbonaceous chondrite OM since it contains ~10-25 wt.% O (11-12).

78

However, determining the O isotope composition of OM is challenging since it tends to be 79 80 intimately mixed with O-rich silicates and oxides at a nano- to micro-scale in carbonaceous chondrites (e.g., 18). Acid-maceration used to isolate the insoluble OM (IOM) fraction from 81 whole rock samples removes most of the silicates but is less effective at dissolving sulfides and 82 some refractory O-bearing oxides such as chromite, spinel, hibonite or corundum. Bulk pyrolysis 83 O isotope analysis of IOM is thus susceptible to contamination by residual mineral inclusions. To 84 85 constrain the triple O isotope composition of carbonaceous chondrite IOM, we integrate here high spatial resolution secondary ion mass spectrometry (SIMS) data obtained using NanoSIMS 86 with high precision ^{17,18}O/¹⁶O isotope ratios obtained using large geometry multi-collector IMS 87 88 1270/80 ion probes (referred as L-SIMS in the following). For each L-SIMS O isotope analysis, measurement of ²⁸Si, ³²S, and ⁵⁶Fe¹⁶O intensities allowed a first order filtering of data for which 89 90 the O signals were largely affected by contamination by residual silicate and/or oxide phases (see 91 Methods for details). The results presented here constitute the first high precision triple O isotope 92 estimates for IOM residues isolated from two emblematic carbonaceous chondrite falls, the 93 Ivuna-type (CI) Orgueil meteorite and the Mighei-type (CM) Murchison meteorite.

94

95 **Results**

The δ^{17} O and δ^{18} O values measured by L-SIMS in the Orgueil, Murchison and Cold Bokkeveld IOM residues range between -23.3 ± 2.4 ‰ and +18.9 ± 2.4 ‰ and -18.0 ± 2.3 ‰ and +16.9 ± 98 2.3 ‰ (uncertainties reported at 2 σ), respectively (Fig. 1 and SI Appendix, Table S1). Least 99 square regression through all the data yields a line defined by $\delta^{17}O = 1.00 (\pm 0.14) \times \delta^{18}O - 3.78$ 100 (± 1.35) (95% confidence level, n = 36, $r^2 = 0.86$), which is indistinguishable from the 101 relationship known as the carbonaceous chondrite anhydrous mineral (CCAM) line ($\delta^{17}O = 0.95$ 102 $\times \delta^{18}O - 4.18$) (Fig. 1).

103

The NanoSIMS data allow determining O isotope ratios with larger uncertainties than those 104 obtained by L-SIMS, but with higher spatial resolution, i.e., over region of interests (ROI) that 105 can be selected from ion imaging to minimize the effect of residual oxides and/or silicates, 106 located through analysis of ²⁸Si and ⁵⁶Fe¹⁶O simultaneously with O isotopes. The NanoSIMS 107 analyses obtained over 40 μ m² areas in Murchison and Orgueil IOM residues show little ²⁸Si 108 hotspots but more abundant micron size ¹⁶O-enriched areas (Figs. 2 and S2). The ¹⁶O hotspots are 109 generally associated with ⁵⁶Fe¹⁶O hotspots, suggesting that magnetite and/or chromite are the 110 main mineral phases that have resisted acid-maceration treatments. 111

112

In terms of surface, these ¹⁶O-enriched ROI represent 16-18 % and 12-21 % of the total 40 μ m × 40 μ m areas analyzed in Murchison and Orgueil, respectively, while pure IOM ROI, defined based on ¹⁶O intensity maps (see Methods), comprise 21-31 % and 18-20 % of the Murchison and Orgueil analyzed areas, respectively (Fig. 3 and SI Appendix, Table S2). The processed NanoSIMS data indicate that the $\delta^{17,18}$ O values of the residual O-rich inclusions in Murchison tend to be *ca*. 20-40 ‰ lower than the $\delta^{17,18}$ O values obtained for O-rich inclusions in Orgueil (SI Appendix, Table S2 and Fig. S3). The $\delta^{17,18}$ O values obtained for Murchison pure IOM areas are around 0 to +10 ‰, while Orgueil pure IOM areas tend to have higher $\delta^{17,18}$ O values around +10

to +30 ‰ (Fig. 1). Overall, the O isotope composition estimated for Murchison and Orgueil pure

122 IOM using NanoSIMS are roughly consistent with the ^{17,18}O-rich end of the trends defined by

- 123 Murchison and Orgueil acid-residues L-SIMS data (Fig. 1).
- 124

125 **Discussion**

126 Assessing the level of contamination from residuals micro-inclusions in IOM

127 The main challenge in determining the O isotope composition of IOM isolated from

128 carbonaceous chondrites is related to the presence of residual nano- to micro-inclusions that have

resisted acid-maceration, as shown here by NanoSIMS imaging. The consistency between O

130 isotope values estimated for pure IOM based on high resolution NanoSIMS analyses and the

131 most ^{17,18}O-rich compositions obtained by L-SIMS for Murchison and Orgueil acid-residues,

respectively, suggest that the latter provide an accurate estimate for the O isotope composition of

133 Murchison and Orgueil IOM. We thus consider here that the average values calculated from the

two most ^{17,18}O-rich compositions measured by L-SIMS on both Murchison and Orgueil provide

us with the best estimates for the O isotope compositions of pure IOM end-members in these

136 samples. This yields $\delta^{18}O = +4.7 \pm 7.7 \%$ (2SD) and $\delta^{17}O = +2.9 \pm 10.3 \%$ (2SD) for Murchison

137 IOM and
$$\delta^{18}O = +16.6 \pm 0.8 \text{ (2SD)}$$
 and $\delta^{17}O = +17.0 \pm 5.2 \text{ (2SD)}$ for Orgueil IOM

138

Figure 4 presents the results of mixing calculations, where O isotope compositions have been calculated for mixed compositions between the IOM $\delta^{17,18}$ O values calculated from the ^{17,18}O-rich L-SIMS data for Murchison and Orgueil and the most negative $\delta^{17,18}$ O values measured in mineral phases in both Murchison (*ca.* -40‰ for spinel; 21) and Orgueil (*ca.* -10‰ for olivine;

143	22). These calculations show that up to \sim 50 % and \sim 80 % contamination of the Murchison and
144	Orgueil O signals, respectively, by ¹⁶ O-rich residual mineral phases could explain the spread of
145	$\delta^{17,18}$ O values measured by L-SIMS (Fig. 4). Such levels of contamination are consistent with
146	NanoSIMS data obtained over 40 μm^2 areas selected randomly in Murchison and Orgueil acid-
147	residues (SI Appendix, Fig. S3). In addition, these calculations are also consistent with O-rich
148	contaminants in Murchison being characterized by $\delta^{17,18}$ O values <i>ca</i> . 20-40 ‰ lower than those
149	of O-rich inclusions in Orgueil (SI Appendix, Table S2 and Fig. S3), even though it is not
150	possible to estimate their true $\delta^{17,18}$ O values corrected for NanoSIMS instrumental mass
151	fractionation since their exact mineralogy was not determined.

152

153 <u>Comparison of bulk and SIMS-derived oxygen isotope data</u>

Because O-rich residual contaminants have lower $\delta^{17,18}$ O values than IOM, it is important to note 154 that the pure IOM O isotope compositions calculated from ^{17,18}O-rich L-SIMS analyses provide 155 minimum estimates. The δ^{18} O and δ^{17} O values estimated for Orgueil IOM by L-SIMS (+16.6 ± 156 0.8 ‰ and +17.0 ± 5.2 ‰, respectively) are higher than the bulk δ^{18} O and δ^{17} O values 157 determined by Halbout et al. (23) (+6.0 \pm 0.8 ‰ and +3.4 \pm 0.4 ‰, respectively), which may 158 indicate that their O isotope ratios were also affected by contamination issues. On the other hand 159 the δ^{18} O value estimated for Orgueil IOM by L-SIMS is in good agreement with the bulk δ^{18} O 160 value of +14.5 \pm 0.6 % determined by Alexander et al. (11). For Murchison IOM, the δ^{18} O value 161 calculated from L-SIMS analyses of +4.7 \pm 7.7 ‰ (2SD) is lower than the bulk IOM δ^{18} O values 162 of $+13.8 \pm 1.6$ % (n = 2, 2SD) determined by Alexander et al. (11-12). These authors take 163 164 advantage of the oxidation of sulfides in air over several days to gradually remove them from

165	acid-residues (11). Modification of the O isotope composition of organic O-bearing chemical
166	functions during such oxidation of acid-residues in air may be a possibility to explain the
167	discrepancy between bulk and L-SIMS δ^{18} O values obtained for Murchison IOM, since
168	atmospheric O ₂ has a δ^{18} O value of <i>ca</i> . +23-24 ‰ (24-25). It is also possible that our L-SIMS
169	δ^{18} O estimate for Murchison IOM does not correspond to pure IOM but to IOM still
170	contaminated by minute amounts of O-rich residual inclusions. At this stage it is not possible to
171	favor one of these two hypotheses over the other. Interestingly, δ^{18} O values obtained on CM
172	chondrite bulk IOM are characterized by much larger variations (from -3.7 ‰ in Essebi to $+14.4$
173	‰ in Murchison, both meteorites being observed falls) compared to other chondrite types (11-
174	12), which may either be related to variable contamination issues of bulk analyses or indicate that
175	CM chondrites accreted IOM with variable O isotope compositions. Clearly, further bulk and in
176	situ investigations are required to fully explore this issue.

177

178 *Triple oxygen isotope constraints on the origin of carbonaceous chondrite IOM*

The O isotope compositions estimated for Murchison IOM ($\delta^{17,18}$ O = *ca*. +3-5 ‰) and Orgueil 179 IOM ($\delta^{17,18}$ O = ca. +17 ‰) fall on the slope 1 line in a δ^{17} O vs. δ^{18} O diagram (Fig. 5). As 180 highlighted by Alexander et al. (11), δ^{18} O values of the CI and CM chondrite IOM are similar to 181 those of their matrix component (Fig. 5). However, the bulk and matrix δ^{17} O values obtained in 182 CI chondrites appear to be lower compared to the δ^{17} O of Orgueil IOM (Fig. 5). These new 183 results, combined with the O isotope composition of CI-CM chondrite original anhydrous 184 silicates, of matrix silicates, and of primordial alteration waters (which are thought to be similar 185 for CI and CM chondrite parent bodies; 21, 26-29), thus seem to rule out scenarios in which the 186

187 O isotope composition of O-bearing functional groups in CI chondrite IOM resulted from O 188 isotope exchange between organic precursors and silicate components during aqueous alteration 189 on the CI chondrite parent body (Fig. 5). On the other hand, because of the larger uncertainty 190 associated with the $\delta^{17,18}$ O estimates for Murchison IOM, its formation during hydrothermal 191 alteration on the CM parent body cannot be totally excluded.

192

The limited existing O isotope dataset obtained on a handful of carbonaceous chondrite 193 meteorites so far suggest that IOM in the CI chondrite Orgueil tends to be enriched in ^{17,18}O 194 compared to IOM in the CM chondrite Murchison (Fig. 5). Interestingly, this relationship is 195 consistent with the variations of average H and N isotope compositions in CM and CI IOM, 196 where CI IOM is enriched in D and ¹⁵N compared to CM IOM (11-12). The D and ¹⁵N 197 enrichments commonly observed in CC IOM have generally been attributed to low temperature 198 processes (<150 K) such as ion-molecule reactions taking place in dense interstellar media or at 199 200 the surface of the PSN. However, recent experimental studies focused on the IOM molecular structure (9, 39), its bulk D/H (40) and the occurrence of D/H hotspots (41), and its noble gas 201 202 isotope signatures (9), have argued that CC IOM could be produced by photochemical reactions 203 involving organic radicals and taking place in the PSN regions where solar UV irradiation would 204 have induced dissociation of $C_x H_y$ molecules. Experiments have shown that photochemical 205 reactions can produce mass-independent O isotope anomalies (e.g., 42-44). One could thus 206 postulate that the mass-independent isotopic fractionation of oxygen isotopes in CC IOM also 207 resulted from chemical reactions involving radical chemistry of CHON-bearing species in the PSN. If correct, such an effect now remains to be experimentally documented in a setting relevant 208 209 to organics formation.

210

211 Cosmochemical implications

The O isotope compositions estimated for CI-CM chondrite IOM fall on a slope 1 line in a δ^{17} O 212 vs. δ^{18} O diagram, which, at a larger scale, describes the O isotope variations of most Solar 213 System objects such as the Sun, high temperature phases (i.e., CAI and chondrules) formed 214 215 during the first few million years of Solar System evolution, and terrestrial planets for example (Fig. 5). Yet, the origin of this slope 1 line in planetary materials is still unknown. A possible 216 mechanism involves self-shielding of ¹⁶O-rich CO gas by UV light during photo-dissociation 217 (e.g., 42), but whether this occurred in the presolar molecular cloud (45) or in the PSN (46) 218 219 remains debated. O isotope compositions of CC IOM appear to fall on a slope 1 line; it could thus be argued that oxygen contained in CC IOM derived from a combination of that found in ¹⁶O-rich 220 CO and ^{17,18}O-rich H₂O molecules formed as a result of self-shielding. If the different CC parent 221 222 bodies accreted IOM sourced from a common carbonaceous reservoir, formed in the presolar molecular cloud, one may expect the various asteroidal parent bodies to have accreted presolar 223 224 IOM grains characterized by similar O isotope compositions. The different triple O isotope 225 compositions for Orgueil and Murchison IOM, dispersed along the slope 1 line, do not seem to 226 favor such a scenario. Alternatively, and considering that Murchison and Orgueil IOM O isotope compositions are representative of those of CM and CI IOM in general, the observation that 227 $\delta^{17,18}O_{CM IOM} < \delta^{17,18}O_{CI IOM}$, consistently with what has been measured for H and N isotope data 228 $(\delta D_{CM IOM} < \delta D_{CI IOM} \text{ and } \delta^{15} N_{CM IOM} < \delta^{15} N_{CI IOM}; 11-12)$, may indicate that carbonaceous 229 230 asteroids accreted IOM that formed locally in the PSN through photochemical radical chemistry involving CHON-bearing species (9, 41). Because of its elevated δD and $\delta^{15}N$ values, it has been 231 proposed that CR IOM could represent the least processed IOM component accreted in 232

carbonaceous asteroids (see discussion in ref. 2). Based on the observed relationship between H, N and O isotope compositions in CI and CM chondrite IOM, we would expect CR chondrite IOM to have $\delta^{17,18}$ O values higher than those of CI chondrite IOM. Determining with high precision the triple O isotope composition of CR chondrite IOM would thus provide important constraints to further explore the formation mechanism(s) of carbonaceous chondrite IOM.

238

239 Materials and Methods

240 <u>Organic matter isolation</u>

IOM was isolated from the Orgueil, Murchison, and Cold Bokkeveld carbonaceous chondrite 241 242 meteorites through successive demineralization using a HF-HCl acidic treatment (47). Powdered meteorite samples were first stirred at room temperature in water, followed by CH₂Cl₂/MeOH 243 244 (2/1, v/v), in order to remove soluble organic compounds. Carbonates were then removed at room temperature using HCl 6N to minimize the formation of fluorides during HF/HCl maceration. 245 246 Samples were then centrifuged and washed with distilled water until reaching neutrality. Isolation of IOM was achieved through acid maceration at room temperature in a HF/HCl mixture (2/1, 247 v/v). Samples were further centrifuged and washed with distilled water to reach neutrality. 248 Neoformed fluorides were then degraded using HCl 6N at 60°C for 24 hours. After HCl hot acid 249 250 maceration, IOM residues were washed with distilled water until reaching neutrality and 251 thoroughly dried. For secondary ion mass spectrometry (SIMS) investigations, a few mg of IOM samples were pressed into high purity indium (99.999 %) and carbon coated. 252

253

254 IMS 1270/80 secondary ion mass spectrometry

Triple O isotope compositions of the IOM samples were measured using the CAMECA IMS 1270 E7 and 1280 HR2 ion probe instruments at the Centre de Recherches Pétrographiques et Géochimiques (CRPG) in Nancy (France) over several analytical sessions, using identical analytical protocols. Negative ${}^{16}O^{-}$, ${}^{17}O^{-}$ and ${}^{18}O^{-}$ secondary ions produced using a ~10 nA Cs⁺ primary beam, accelerated at 10 kV and rastered over ~20 µm diameter areas, were measured in multicollection mode with one Faraday cup (FC) on the L'2 trolley for ${}^{16}O^{-}$ and two electron

multipliers (EM) for ¹⁷O⁻ (central EM) and ¹⁸O⁻ (H2 EM). To maximize peak flatness, entrance 261 and exit slits were adjusted to achieve a mass resolving power of ~ 8000 for $^{17}O^{-}$ on the central 262 EM and ~2500 on the off-axis L'2 FC and H2 EM (using slit #1 of the off-axis collectors). 263 Organic matter samples contain significant amounts of OH (average ${}^{16}\text{OH}^{-}/{}^{17}\text{O}^{-} \sim 97 \pm 17$, ~120 ± 264 31 and \sim 121 \pm 20 in Orgueil IOM, Murchison IOM and Cold Bokkeveld IOM, respectively) and 265 the protocol used did not completely eliminate contribution from the ${}^{16}OH^{-1}$ tail on the ${}^{17}O^{-1}$ peaks. 266 To quantify this contribution and adequately correct the measured ${}^{17}O/{}^{16}O$ ratios, we assumed 267 that the ¹⁶OH⁻ peak was symmetrical, calculated the mass difference between the center of the 268 ${}^{17}\text{O}^{-}$ (16.9991 amu) and ${}^{16}\text{OH}^{-}$ (17.0027 amu) peaks and counted the ${}^{16}\text{OH}^{-}$ tail intensity at mass 269 17.0063 amu (mass ${}^{16}\text{OH}^-$ + [mass ${}^{16}\text{OH}^-$ - mass ${}^{17}\text{O}^-$]) for 50 s before and after each analysis. The 270 $^{16}\text{OH}^-$ tail/peak ratios were ~1.5 \pm 0.5 \times 10 $^{-5}$ in Murchison IOM, ~1.1 \pm 0.1 \times 10 $^{-5}$ in Cold 271 Bokkeveld and $\sim 1.9 \pm 0.5 \times 10^{-5}$ in Orgueil IOM. This resulted in correction of the measured 272 δ^{17} O values by 0.7-2.8 ‰ in Murchison IOM, 1.3-1.8 ‰ in Cold Bokkeveld IOM and 0.8-2.4 ‰ 273 in Orgueil IOM (SI Appendix, Table S1). For each analysis, the FC background was measured 274 during pre-sputtering. Deadtime of the EM was also calibrated once per analytical session. The 275 276 total analysis time was 260 s (60 s pre-sputtering and 40 cycles of 5 s each measurement time). Instrumental mass fractionation (IMF) for O isotope measurements in IOM samples was 277 corrected by repeated analyses of our Clarno kerogen standard ($\delta^{18}O_{bulk} = 14.3 \pm 0.1$ ‰; 48), for 278 which we assumed a $\delta^{17}O_{\text{bulk}}$ of 7.4 ‰, i.e., a terrestrial O isotope composition. Count rates 279 obtained on the Clarno kerogen standard were $0.3-1.2 \times 10^7$ cps nA⁻¹ for ¹⁶O⁻, $1.1-4.5 \times 10^3$ cps 280 nA^{-1} for ${}^{17}O^{-}$ and $0.6-2.3 \times 10^4$ cps nA^{-1} for ${}^{18}O^{-}$, similar to those obtained on the IOM samples 281 $(0.3-3.1 \times 10^7 \text{ cps nA}^{-1} \text{ for } {}^{16}\text{O}^{-}, 0.1-1.1 \times 10^4 \text{ cps nA}^{-1} \text{ for } {}^{17}\text{O}^{-} \text{ and } 0.5-5.5 \times 10^4 \text{ cps nA}^{-1} \text{ for } {}^{18}\text{O}^{-}$ 282). The final uncertainties for individual $\delta^{17,18}$ O values, reported in SI Appendix (SI Appendix, 283 284 Table S1) at the 2σ level, include uncertainties related to counting statistics associated with each individual analysis and the external reproducibility measured for $\delta^{17,18}$ O values on the Clarno 285 kerogen standard. Over three analytical sessions in February 2016, July 2016 and December 286 2016, we obtained a weighted average Δ^{17} O of -0.1 ± 0.4 ‰ (95% confidence level, n = 66, 287 MSWD = 3.0) (SI Appendix, Fig. S1). We further tested our analytical protocol on the Silurian 288 Zdanow terrestrial kerogen and obtained an average $\delta^{18}O_{SIMS}$ of $12.4 \pm 4.6 \% (2SD - n = 9)$, 289 which is consistent with its bulk δ^{18} O of 13.4 ± 0.2 ‰ (48). The Δ^{17} O measured on Zdanow was 290

- 291 $0.2 \pm 1.8 \% (2SD n = 9)$, indicating that Zdanow sits on the TFL, which shows that our L-292 SIMS protocol accurately measures the triple O isotope composition of organic residues.
- 293

The secondary species ${}^{12}C^{1}H$, ${}^{16}O$, ${}^{28}Si$, ${}^{32}S$ and ${}^{56}Fe^{16}O$ were collected following O isotope analyses on the same analytical spots using the magnetic peak switching mode and a ~10 nA Cs⁺ beam in order to identify and filter the IOM data largely affected by contamination by residual silicate and oxide phases (see details in refs. 48-49).

298

299 <u>Nanoscale secondary ion mass spectrometry</u>

The triple O isotope composition of the Orgueil and Murchison IOM residues was also measured 300 using the CAMECA NanoSIMS 50L ion probe instrument at The University of Manchester 301 (UK). Negative ${}^{16}O^{-}$, ${}^{17}O^{-}$, ${}^{18}O^{-}$, ${}^{12}C_2^{-}$, ${}^{12}C_2^{-14}N^{-}$, ${}^{28}Si^{-}$ and ${}^{56}Fe^{16}O^{-}$ secondary ion species produced 302 using a ~15 pA Cs⁺ primary beam, accelerated at 16 kV and rastered over 40 μ m × 40 μ m areas, 303 304 were measured in multicollection mode on seven electron multipliers (EM). Before analysis, a ~100 pA Cs⁺ primary beam was rastered over 50 μ m × 50 μ m areas for 5 minutes to clean the 305 sample surface and reach sputtering equilibrium. To limit the ¹⁶OH⁻ interference on the ¹⁷O⁻ peak, 306 a 10 µm wide entrance slit (ES5) was used at the entrance of the mass analyzer, and a 150 µm 307 wide aperture slit (AS3) reduced the beam divergence, resulting in a mass resolving power of 308 ~8000. An electron gun was used for charge compensation. Using these conditions, the count 309 rates were 15000-35000 cps for 16 O, 500-1800 cps for 12 C₂, 3000-10000 cps for 12 C¹⁴N and 5-20 310 cps for ⁵⁶Fe¹⁶O, ensuring no detector ageing over the week-long analytical session. During the 311 session, the vacuum in the analysis chamber remained constant at ~ 3×10^{-10} mbar. For data 312 acquisition, the 40 μm^2 areas were divided in 256 \times 256 pixels and between 320 and 600 frames 313 314 were acquired at 1000 μ s/px, resulting in a total analysis time of 6 to 11 hours per analysis. Automatic alignment of the secondary beam (EOS, Cy, and P2/P3) and of the peak positions was 315 performed every 50 frames during each analysis based on scanning the ¹⁶O⁻ peak. IMF for O 316 isotope measurements in IOM samples was corrected by analyzing the same Clarno kerogen 317 standard used for the L-SIMS analyses (see above). The final uncertainties for individual $\delta^{17,18}$ O 318 values, reported in SI Appendix (SI Appendix, Table S2) at the 2σ level, include those related to 319 320 counting statistics for each individual analysis and the external reproducibility measured for

321 $\delta^{17,18}$ O on the Clarno kerogen standard (±11.1 ‰ and ±11.9 ‰ for δ^{18} O and δ^{17} O, respectively; 322 2SE, n = 3).

323

324 The NanoSIMS data were processed offline using the l'Image software package (L. Nittler, Carnegie Institution of Washington). A 44 ns deadtime was applied to all EM, and individual 325 frames were binned into packages of 6-10 frames for each analysis to handle these large dataset 326 more easily. Regions of interest (ROI) were defined using lower and upper thresholds for the 327 different species, and comprised, for each analysis, the whole analyzed area, an area with 328 intermediate ¹⁶O intensity and a ¹⁶O-rich area corresponding to hotspots. A ROI of 'pure IOM' 329 was then defined by subtracting the ROI corresponding to the area with intermediate ¹⁶O intensity 330 to the ROI corresponding to the whole analyzed area. The ${}^{17}O/{}^{16}O$, ${}^{18}O/{}^{16}O$, ${}^{12}C^{14}N/{}^{12}C_2$ and 331 $^{16}\text{O}/^{12}\text{C}_2$ ratios were calculated using l'Image. 332 333 All processed data are given in SI Appendix (SI Appendix, Tables S1 and S2). Raw data can be 334 requested to the corresponding author. 335 336 Acknowledgments 337 338 We thank S. Derenne for providing the Murchison IOM sample, N. Bouden and J. Villeneuve for their help with L-SIMS analyses in Nancy, and I.C. Lyon for his help with NanoSIMS analyses 339 340 in Manchester. We also thank the anonymous referees for their constructive reviews. This is IPGP contribution #XXXX and CRPG contribution #XXXX. 341 342 343 344 Funding: This work was supported by the ERC grant PaleoNanoLife (No. 290861; PI F. Robert), 345 the ANR grant CRADLE (ANR-15-CE31-0004-01; PI M. Chaussidon), and the UK Science and Technology Facilities Council grants No. ST/M001253/1 (project Co-I I.C. Lyon) and 346 ST/P005225/1 (PI R. Tartèse). The NanoSIMS at the University of Manchester was funded by 347 348 UK Research Partnership Investment Funding (UKRPIF) Manchester RPIF Round 2. 349

350

351 **References**

- 1. Hayes JM (1967) Organic constituents of meteorites a review. *Geochim Cosmochim Acta* 31:1395-1440.
- Alexander CMO'D, Cody GD, De Gregorio BT, Nittler LR, Stroud RM (2017) The nature,
 origin and modification of insoluble organic matter in chondrites, the major source of Earth's
 C and N. *Chem Erde Geochem* 77:227-256.
- 357 3. Robert F, Epstein S (1982) The concentration and isotopic composition of hydrogen, carbon
 and nitrogen in carbonaceous meteorites. *Geochim Cosmochim Acta* 46:81-95.
- 4. Yang J, Epstein S (1983) Interstellar organic matter in meteorites. *Geochim Cosmochim Acta*47:2199-2216.
- 5. Busemann H, et al. (2006) Interstellar chemistry recorded in organic matter from primitive
 meteorites. *Science* 312:727-730.
- 6. Remusat L, Palhol F, Robert F, Derenne S, France-Lanord C (2006) Enrichment of deuterium
 in insoluble organic matter from primitive meteorites: a solar system origin? *Earth Planet Sci Lett* 243:15-25.
- 366 7. Gourier D, et al. (2008) Extreme deuterium enrichment of organic radicals in the Orgueil
 367 meteorite: revisiting the interstellar interpretation? *Geochim Cosmochim Acta* 72:1914-1923.
- 368 8. Derenne S, Robert F (2010) Model of molecular structure of the insoluble organic matter
 369 isolated from Murchison meteorite. *Meteorit Planet Sci* 45:1461-1475.
- 9. Kuga M, Marty B, Marrocchi Y, Tissandier L (2015) Synthesis of refractory organic matter in
 the ionized gas phase of the solar nebula. *Proc Natl Acad Sci USA* 112:7129-7134.
- 10. Cody GD, et al. (2011) Establishing a molecular relationship between chondritic and
 cometary organic solids. *Proc Natl Acad Sci USA* 108:19171-19176.
- 11. Alexander CMO'D, Fogel M, Yabuta H, Cody GD (2007) The origin and evolution of
 chondrites recorded in the elemental and isotopic compositions of their macromolecular
 organic matter. *Geochim Cosmochim Acta* 71:4380-4403.
- 12. Alexander CMO'D, et al. (2010) Deuterium enrichments in chondritic macromolecular
 material implications for the origin and evolution of organics, water and asteroids. *Geochim Cosmochim Acta* 74:4417-4437.
- 13. Orthous-Daunay FR, et al. (2013) Mid-infrared study of the molecular structure variability of
 insoluble organic matter from primitive chondrites. *Icarus* 223:534-543.
- 14. Quirico E, et al. (2014) Origin of insoluble organic matter in type 1 and 2 chondrites: New
 clues, new questions. *Geochim Cosmochim Acta* 136:80-99.
- 15. Hashiguchi M, Kobayashi S, Yurimoto H (2015) Deuterium- and ¹⁵N-signatures of organic
 globules in Murchison and Northwest Africa 801 meteorites. *Geochem J* 49:377-391.
- 16. Clayton RN (1993) Oxygen isotopes in meteorites. *Ann Rev Earth Planet Sci* 21:115-149.
- 17. McKeegan KD, et al. (2011) The oxygen isotopic composition of the Sun inferred from
 captured solar wind. *Science* 332:1528-1532.

18. Le Guillou C, Bernard S, Brearley AJ, Remusat L (2014) Evolution of organic matter in
 Orgueil, Murchison and Renazzo during parent body aqueous alteration: *In situ* investigations.
 Geochim Cosmochim Acta 131:368-392.

19. Clayton RN, Onuma N, Grossman L, Mayeda TK (1977) Distribution of the presolar
 component in Allende and other carbonaceous chondrites. *Earth Planet Sci Lett* 34:209-224.

- 20. Young ED, Russell SS (1998) Oxygen reservoirs in the early solar nebula inferred from an
- 395 Allende CAI. *Science* 282:452-455.
- 21. Clayton RN, Mayeda TK (1984) The oxygen isotope record in Murchison and other
 carbonaceous chondrites. *Earth Planet Sci Lett* 67:151-161.
- 22. Leshin LA, Rubin AE, McKeegan KD (1997) The oxygen isotopic composition of olivine
 and pyroxene from CI chondrites. *Geochim Cosmochim Acta* 61:835-845.
- 400 23. Halbout J, Robert F, Javoy M (1990) Hydrogen and oxygen isotope compositions in kerogen
 401 from the Orgueil meteorite: Clues to a solar origin. *Geochim Cosmochim Acta* 54:1453-1462.
- 402 24. Kroopnick P, Craig H (1972) Atmospheric oxygen: Isotopic composition and solubility
 403 fractionation. *Science* 175:54-55.
- 404 25. Luz B, Barkan E (2011) The isotopic composition of atmospheric oxygen. *Global*405 *Biogeochem Cycles* 25:GB3001, doi:10.1029/2010GB003883.
- 26. Rowe MW, Clayton RN, Mayeda TK (1994) Oxygen isotopes in separated components of CI
 and CM meteorites. *Geochim Cosmochim Acta* 58:5341-5347.
- 408 27. Clayton RN, Mayeda TK (1999) Oxygen isotope studies of carbonaceous chondrites.
 409 *Geochim Cosmochim Acta* 63:2089-2104.
- 410 28. Fujiya W (2018) Oxygen isotopic ratios of primordial water in carbonaceous chondrites.
 411 *Earth Planet Sci Lett* 481:264-272.
- 412 29. Marrocchi Y, Bekaert DV, Piani L (2018) Origin and abundance of water in carbonaceous
- 413 asteroids. *Earth Planet Sci Lett* 482:23-32.
- 30. Aléon J, El Goresy A, Zinner E (2007) Oxygen isotope heterogeneities in the earliest
 protosolar gas recorded in a meteoritic calcium–aluminum-rich inclusion. *Earth Planet Sci Lett* 263:114-127.
- 417 31. Krot AN, et al. (2008) Oxygen isotopic compositions of Allende Type C CAIs: Evidence for
- 418 isotopic exchange during nebular melting and asteroidal metamorphism. *Geochim Cosmochim*419 *Acta* 72:2534-2555.
- 32. Bodénan JD, Starkey NA, Russell SS, Wright IP, Franchi IA (2014) An oxygen isotope study
 of Wark-Lovering rims on type A CAIs in primitive carbonaceous chondrites. Earth Planet Sci
 Lett 401:327-336.
- 423 33. Clayton RN, et al. (1983) Oxygen isotopic compositions of chondrules in Allende and
- 424 ordinary chondrites. *Chondrules and Their Origins*, ed King EA (Lunar and Planetary
- 425 Institute, Houston), pp 37-43.
- 426 34. Rubin AE, Wasson JT, Clayton RN, Mayeda TK (1990) Oxygen isotopes in chondrules and
- 427 coarse-grained chondrule rims from the Allende meteorite. *Earth Planet Sci Lett* 96:247-255.

35. Weisberg MK, Prinz M, Clayton RN, Mayeda TK (1993) The CR (Renazzo-type) 428 429 carbonaceous chondrite group and its implications. Geochim Cosmochim Acta 57:1567-1586. 430 36. Jones RH, et al. (2004) Oxygen isotope heterogeneity in chondrules from the Mokoia CV3 431 carbonaceous chondrite. Geochim Cosmochim Acta 68:3423-3438. 432 37. Jenniskens P, et al. (2012) Radar-enabled recovery of the Sutter's Mill meteorite, a 433 carbonaceous chondrite regolith breccia. Science 6114:1583-1587. 38. Verdier-Paoletti MJ, et al. (2017) Oxygen isotope constraints on the alteration temperatures 434 435 of CM chondrites. Earth Planet Sci Lett 458:273-281. 39. Biron K, Derenne S, Robert F, Rouzaud JN (2015) Toward an experimental synthesis of the 436 chondritic insoluble organic matter. Meteorit Planet Sci 50:1408-1422. 437 438 40. Laurent B, et al. (2015) The deuterium/hydrogen distribution in chondritic organic matter 439 attests to early ionizing irradiation. Nat Comm 6:8567, doi:10.1038/ncomms9567. 440 41. Robert F, et al. (2017) Hydrogen isotope fractionation in methane plasma. Proc Natl Acad Sci USA 114:870-874. 441 442 42. Thiemens MH, Heidenreich JE III (1983) The mass-independent fractionation of oxygen - a novel isotope effect and its possible cosmochemical implications. Science 219:1073-1075. 443 43. Chakraborty S, Ahmed M, Jackson TL, Thiemens MH (2008) Experimental test of self-444 shielding in vacuum ultraviolet photodissociation of CO. Science 321:1328-1331. 445 44. Chakraborty S, Yanchulova P, Thiemens MH (2013) Mass-independent oxygen isotopic 446 447 partitioning during gas-phase SiO₂ formation. Science 342:463-466. 448 45. Yurimoto H, Kuramoto K (2004) Molecular cloud origin for the oxygen isotope 449 heterogeneity in the solar system. Science 305:1763-1766. 450 46. Lyons JR, Young ED (2005) CO self-shielding as the origin of oxygen isotope anomalies in 451 the early solar nebula. Nature 435:317-320. 452 47. Durand B, Nicaise G (1980) Procedures for kerogen isolation. Kerogen – Insoluble organic matter from sedimentary rocks, ed Durand B (Technip, Paris), pp 35-54. 453 454 48. Tartèse R, Chaussidon M, Gurenko A, Delarue F, Robert F (2016) In situ oxygen isotope 455 analysis of fossil organic matter. Geochim Cosmochim Acta 182:24-39. 49. Tartèse R, Chaussidon M, Gurenko A, Delarue F, Robert F (2017) Warm Archean oceans 456 457 reconstructed from oxygen isotope composition of early-life remnants. Geochem Perspect Lett 3:55-65. 458 459 460 461 462 463

464 **Figure captions**

465

Figure 1: $\delta^{17,18}$ O values obtained in Orgueil, Murchison and Cold Bokkeveld IOM residues. The terrestrial fractionation line (TFL), the carbonaceous chondrite anhydrous mineral line (CCAM; 19) and the Young and Russell line (Y&R; 20) are also represented.

468 I

470 Figure 2: NanoSIMS images showing the distribution of ${}^{12}C^{14}N$, ${}^{16}O$, ${}^{56}Fe^{16}O$ and ${}^{28}Si$ secondary

ion species in Murchison and Orgueil IOM acid-maceration residues. White and cyan arrows

472 indicate higher 56 Fe 16 O and 28 Si intensities, respectively.

473

474 Figure 3: Examples of the regions of interest defined on ¹⁶O ion images for one analysis each of

- 475 Murchison and Orgueil acid-residues. ROI#3 (left) correspond to intermediate O intensity,
- 476 ROI#4 (center) correspond to O hotspots, and ROI#5 (right) correspond to pure IOM (see text for
- 477 details).
- 478

Figure 4: Calculated O isotope values for mixed compositions between the estimated IOM $\delta^{17,18}$ O

480 values (grey stars) in (A) Murchison and (B) Orgueil acid-residues and the most negative $\delta^{17,18}$ O

values measured in mineral phases in both Murchison (ca. -40‰ for spinel; 21) and Orgueil (ca. -

482 10‰ for olivine; 22) (dark grey hexagons). White dots represent 10% mixing intervals, and 20%

483 mixing intervals are given on the diagrams from 0% to 100% mineral contribution.

484

485 Figure 5: O isotope compositions measured by L-SIMS on IOM residues isolated from the

486 Orgueil and Murchison carbonaceous chondrites. The terrestrial fractionation line (TFL), the

487 CCAM line, and the Y&R line, are also represented, together with the O isotope compositions of

the Sun (17), of CI and CM chondrite components (bulk, matrix and anhydrous silicates; 21-22,

489 26-27), of carbonaceous chondrite Ca- and Al-rich inclusions (30-32), chondrules (33-37) and

490 carbonates (38). Estimates for the O isotope composition of CM chondrite primordial water

491 (HW1 & HW2) are from ref. 27 for HW1 and ref. 38 for HW2.

492













Supplementary Information for

Insights into the origin of carbonaceous chondrite organics from their triple oxygen isotope composition

R. Tartèse, M. Chaussidon, A. Gurenko, F. Delarue, F. Robert

Corresponding author: Romain Tartèse Email: <u>romain.tartese@manchester.ac.uk</u>

This PDF file includes:

Figs. S1 to S3 Tables S1 to S2



Fig. S1. Histogram and probability distribution of Δ^{17} O values measured on the Clarno kerogen standard over all the L-SIMS analytical sessions.



Fig. S2. NanoSIMS images obtained on $40 \times 40 \,\mu\text{m}$ areas in Murchison and Orgueil IOM showing the distribution of ${}^{12}\text{C}{}^{14}\text{N}$, ${}^{12}\text{C}{}_2$, ${}^{16}\text{O}$, ${}^{56}\text{Fe}{}^{16}\text{O}$ and ${}^{28}\text{Si}$ secondary ion species.



Fig. S3. Plot showing the NanoSIMS ¹⁶O intensity versus the measured δ^{18} O values for regions of interest corresponding to areas with a mixture of IOM and O-rich residual minerals (ROI3), O-rich hotspots (ROI4), and 'pure IOM' areas (ROI5) for five areas analyzed in Murchison and Orgueil acid-residues.

Analysis	Secon (cps nA	dary ion int ⁻¹ primary o	ensity current)	¹⁶ OH		Isotope ratios (‰ vs. SMOW)							
1 xiidi y 515	¹⁶ O ⁻	¹⁷ O ⁻	¹⁸ O ⁻	(‰)*	δ ¹⁸ Ο	2σ	$\delta^{17}O$	2σ	$\Delta^{17}O$	2σ			
Orgueil IOM													
Orgueil_3iso@4	3.80E+06	1.40E+03	7.21E+03	1.7	-4.2	2.9	-6.2	2.0	-4.0	2.3			
Orgueil_3iso@5	3.98E+06	1.47E+03	7.57E+03	1.3	-1.6	2.9	-5.5	1.9	-4.7	2.1			
Orgueil_3iso@7	4.07E+06	1.50E+03	7.75E+03	0.8	1.0	2.9	-6.9	1.9	-7.4	1.9			
Orgueil@2	3.02E+06	1.13E+03	5.82E+03	2.4	5.9	2.3	2.3	2.4	-0.8	2.2			
Orgueil@3	2.56E+06	9.59E+02	4.94E+03	2.3	6.5	2.3	3.1	2.5	-0.3	2.5			
Orgueil@4	2.33E+06	8.72E+02	4.49E+03	2.3	6.0	2.3	2.8	2.5	-0.3	2.5			
Orgueil@5	6.27E+06	2.37E+03	1.21E+04	1.5	10.9	2.3	10.5	2.5	4.9	2.4			
Orgueil@6	7.54E+06	2.87E+03	1.46E+04	1.5	16.3	2.3	18.9	2.4	10.4	2.2			
Orgueil@9	4.10E+06	1.55E+03	7.99E+03	2.2	16.9	2.3	15.2	2.4	6.4	2.3			
Orgueil@11	2.64E+06	9.92E+02	5.12E+03	2.0	11.8	2.3	6.1	2.5	0.0	2.5			
Orgueil@12	4.28E+06	1.59E+03	8.22E+03	1.3	2.5	2.3	-4.2	2.4	-5.5	2.2			
Orgueil@13	3.64E+06	1.37E+03	7.03E+03	1.4	8.2	2.3	4.2	2.4	-0.1	2.2			
Orgueil@16	2.44E+06	9.12E+02	4.71E+03	1.8	6.5	2.3	0.3	2.5	-3.1	2.4			
Orgueil@17	2.61E+06	9.79E+02	5.08E+03	1.7	15.2	2.3	5.8	2.4	-2.1	2.3			
Orgueil@19	1 50E+07	5 57E+03	2.84E+04	2.1	7.3	2.3	-2.0	2.4	-5.7	2.3			
orgaon e ry	11002107	01072100	Murchis	on IOM									
Murchison 2ice@1	2.57E+07	0.40E+02	4 71E+04	1 7	65	20	10.1	1.0	67	1.7			
Murchison_31so@1	2.37E+07	9.40E+05	4./1E+04	1.7	-0.5	2.9	-10.1	1.9	-0.7	1.7			
Murchison_3iso@2	4.42E+07	1.01E+04	8.14E+04	0.7	-7.5	2.9	-10.1	1.9	-12.2	1.0			
Murchison_Siso@S	2.55E+07	9.45E+05	4.81E+04	1.1	1.9	2.9	-0.7	1.9	-1.7	2.0			
Murchison_3iso@4	9.77E+06	3.66E+03	1.86E+04	1.9	1.4	2.9	0.0	1.9	2.7	1.7			
Murchison_3iso@5	3.05E+07	1.11E+04	5.53E+04	1.1	-13.4	2.9	-1/.8	1.9	-10.9	1.7			
Murchison_3iso@/	1.36E+07	4.96E+03	2.51E+04	1.5	-14.4	3.0	-14.7	2.0	-1.2	2.5			
Murchison_3iso@8	1.55E+07	5.70E+03	2.89E+04	1.0	-4.6	2.9	-/.3	1.9	-4.9	1.8			
Murchison@1	1.66E+07	6.10E+03	3.10E+04	2.8	-4./	2.3	-8.8	2.4	-6.3	2.2			
Murchison@2	1.42E+07	5.17E+03	2.64E+04	2.8	-18.0	2.3	-23.3	2.4	-14.0	2.2			
Murchison@5	2.06E+07	7.51E+03	3.82E+04	1.9	-11.8	2.3	-18.6	2.4	-12.5	2.0			
Murchison@6	1.34E+07	4.92E+03	2.52E+04	2.5	-9.3	2.3	-17.2	2.4	-12.4	2.1			
Murchison@8	7.39E+06	2.73E+03	1.40E+04	4.9	-10.3	2.3	-14.5	2.5	-9.1	2.4			
Murchison@9	1.73E+07	6.31E+03	3.21E+04	2.0	-14.3	2.3	-19.7	2.4	-12.2	2.2			
			Cold Bokk	eveld IO	М								
CB_3iso@1	1.41E+07	5.12E+03	2.61E+04	1.5	-13.1	2.9	-19.8	2.0	-13.0	2.3			
CB_3iso@2	1.56E+07	5.70E+03	2.90E+04	1.4	-9.4	2.9	-14.9	2.0	-10.0	2.2			
CB_3iso@3	1.11E+07	4.07E+03	2.06E+04	1.6	-14.5	2.9	-8.8	1.9	-1.3	2.0			
CB_3iso@4	1.35E+07	4.91E+03	2.52E+04	1.3	-10.9	2.9	-23.0	1.9	-17.3	1.9			
CB_3iso@5	1.20E+07	4.42E+03	2.24E+04	1.3	-9.8	2.9	-10.9	1.9	-5.8	1.7			
CB_3iso@6	1.52E+07	5.59E+03	2.83E+04	1.3	-5.9	2.9	-5.6	1.9	-2.5	1.6			
CB_3iso@7	8.59E+06	3.16E+03	1.61E+04	1.8	-10.0	2.9	-12.1	1.9	-6.9	1.9			
CB 3iso@8	1.71E+07	6.32E+03	3.19E+04	1.1	-5.2	2.9	-4.7	1.9	-2.0	1.8			

Table S1. L-SIMS oxygen isotope results. ^{*}Indicates the contribution of the ${}^{16}OH^{-}$ peak tail on the measured ${}^{17}O^{-}$ intensity.

DOI	Area	Intensity (cps)						Ratio (in ‰ vs. SMOW for $\delta^{17,18}$ O)								IN	IMF corrected (%)			
KUI	(µm ²)	¹⁶ O	¹⁷ O	¹⁸ O	¹² C ₂	¹² C ¹⁴ N	²⁸ Si	⁵⁶ Fe ¹⁶ O	δ ¹⁸ Ο	1σ	$\delta^{17}O$	1σ	$^{12}C^{14}N/^{12}C_2$	1σ	¹⁶ O/ ¹² C ₂	1σ	δ ¹⁸ Ο	2σ	$\delta^{17}O$	2σ
Murchison #1																				
All area	1517	31880	11.6	59.7	1607.5	6399.7	0.9	36.0	-65.7	0.6	-50.6	1.5	3.88	0.04	19.8	0.2				
Broad O rich	1091	38251	13.9	71.6	1704.2	6710.0	1.1	45.2	-66.8	0.7	-51.7	1.6	3.84	0.04	22.4	0.3				
O hotspot	270	68513	24.7	127.7	1591.3	6259.3	1.1	94.6	-70.9	1.0	-58.6	2.3	3.84	0.04	43.0	0.5				
'pure' IOM	314	18985	6.9	35.8	1736.2	6985.5	0.9	14.5	-58.8	1.9	-50.2	4.4	3.92	0.04	10.8	0.2	-0.3	11.7	-8.1	14.8
Murchison #2																				
All area	1529	39379	14.5	74.5	1669.1	7047.1	1.8	40.7	-56.6	0.7	-40.5	1.7	4.10	0.05	23.1	0.4				
Broad O rich	1191	47012	17.2	89.0	1772.4	7463.7	2.0	50.8	-56.2	0.8	-40.7	1.7	4.10	0.05	26.0	0.4				
O hotspot	268	92795	33.9	175.7	1688.2	7168.4	3.1	123.6	-55.8	1.1	-46.0	2.6	4.14	0.04	52.9	1.0				
'pure' IOM	475	21818	8.1	41.4	1833.6	7867.2	1.3	15.9	-54.7	1.8	-31.2	4.1	4.17	0.05	11.6	0.2	3.8	11.7	10.9	14.4
Murchison #3																				
All area	1548	38520	14.1	72.4	1550.8	7138.4	1.0	23.8	-62.6	1.0	-44.6	1.7	4.44	0.06	24.7	0.4				
Broad O rich	999	51440	18.8	96.5	1622.3	7371.6	1.1	32.2	-64.5	1.0	-45.9	1.9	4.39	0.05	31.6	0.4				
O hotspot	249	109608	39.7	204.4	1445.1	6530.8	1.0	64.9	-70.4	1.5	-53.0	2.5	4.37	0.05	76.4	0.7				
'pure' IOM	400	16439	6.1	31.3	1640.8	7863.2	0.8	9.7	-51.4	2.2	-36.7	5.2	4.64	0.05	9.5	0.2	7.1	12.0	5.4	15.8
Orgueil #3																				
All area	1548	27211	10.1	52.8	816.5	3682.9	1.3	9.8	-32.5	0.9	-35.0	2.0	4.27	0.07	33.2	0.4				
Broad O rich	745	35271	13.0	68.4	756.8	3574.5	1.5	13.2	-32.1	1.1	-34.2	2.5	4.47	0.07	46.4	0.6				
O hotspot	191	49642	18.3	96.0	672.7	3345.4	1.7	19.9	-35.6	1.8	-44.4	4.1	4.72	0.07	73.2	1.1				
'pure' IOM	314	17747	6.6	34.3	946.2	4072.3	1.1	5.4	-36.6	2.4	-34.9	5.6	4.08	0.07	18.6	0.2	21.9	12.2	7.2	16.3
Orgueil #4																				
All area	1531	23537	8.9	46.1	712.6	3273.8	2.0	15.3	-22.8	1.0	-12.3	2.3	4.26	0.09	32.1	0.6				
Broad O rich	901	32093	12.2	62.9	661.0	3333.3	2.9	23.1	-21.6	1.1	-12.2	2.5	4.65	0.11	48.0	0.8				
O hotspot	323	49644	18.9	97.8	465.2	2801.3	5.4	42.5	-18.4	1.5	-3.2	3.5	5.53	0.13	106.5	1.6				
'pure' IOM	272	12251	4.6	23.8	993.2	4146.7	0.7	3.8	-30.5	3.3	-19.5	7.5	3.89	0.08	11.5	0.3	28.0	13.0	22.6	19.2

 Table S2. NanoSIMS results obtained on Murchison and Orgueil acid-maceration residues.