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1	Thermal recalcitrance of the organic D-rich component of ordinary
2	chondrites
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10	
11	ABSTRACT
12	
13	Carbonaceous and ordinary chondrites (CCs and OCs) contain insoluble organic
14	matter (IOM) with large D-excess compared to other objects in the solar system. The higher
15	the temperature experienced by CCs, the lower the D/H ratio of their IOM. It seems to be the
16	opposite for OCs. Here, we report NanoSIMS H- (and N-) isotopic imaging of IOM of three
17	OCs that experienced thermal metamorphism in the sequence Semarkona, Bishunpur and
18	GRO 95502. In addition, we performed flash heating experiments on the IOM of GRO 95502
19	at 600°C and characterized the residues using NanoSIMS, Raman and XANES spectroscopy.
20	The present study shows that, in contrast to IOM of CI, CM and CR, IOM of OCs exhibits
21	very few D-rich (or ¹⁵ N-rich) hotspots. Furthermore, although the evolution of the molecular
22	structure of OC and CC IOM is similar upon heating, their D/H ratios do not follow the same
23	trend: the D/H of OC IOM drastically increases while the D/H of CC IOM decreases. In
24	contrast to CC IOM, the D-rich component of which does not survive at high temperatures,
25	the present results highlight the thermal recalcitrance of the D-rich component of OC IOM.

This suggests that CCs and OCs did not accrete the same organic material, thereby challenging the hypothesis of a common precursor on chondritic parent bodies. The present results support the hypothesis that OC IOM contains an organic component that could originate from the interstellar medium.

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Keywords: insoluble organic matter; chondrite; parent body processes; NanoSIMS; hydrogen
and nitrogen isotopes; solar system

1. INTRODUCTION

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36 Chondrites are often considered to be the witnesses of solar system formation (Weisberg et al., 2006). Their components are believed to have formed during the collapse of 37 38 the parent molecular cloud and solar nebula evolution. They are consequently the targets for 39 many studies aiming to understand the processes of formation of the constituents of 40 planetesimals and planets. The origin of volatile reservoirs in the early solar system is the 41 subject of intense debate, with implications for the origin of water and life on Earth. In this 42 respect, carbonaceous chondrites (CCs) have received most of the attention because they are 43 the most organic- and water-rich chondrites (CCs may contain up to a few weight percent of 44 carbon). Two types of organic matter are usually distinguished (Remusat, 2015): the organic 45 matter that is soluble in water and organic solvents, and the insoluble organic matter (IOM). 46 The IOM constitutes the major part of the total organic material and is generally studied after 47 isolation from the minerals by acid treatments using HF-HCl (Robert and Epstein, 1982) or 48 CsF-HCl, (Alexander et al., 2007).

49 The molecular, structural and chemical composition of CC IOM has been extensively 50 documented (Remusat, 2015 and references herein). It corresponds to a macromolecule made 51 of small aromatic units linked by short aliphatic chains that contains hetero-elements (N, S, 52 O) (Havatsu et al., 1980; Remusat et al., 2005; Cody and Alexander, 2005; Orthous-Daunav et al., 2010) and organic radicals (Binet et al., 2002; Binet et al., 2004). Type 1 and 2 CC IOM 53 54 is enriched in heavy isotopes of H and N relative to terrestrial organic matter, with the 55 exception of some CMs having near-atmospheric IOM N isotopic compositions (Robert and 56 Epstein, 1982; Yang and Epstein, 1983; Alexander et al., 2007; Alexander et al., 2010), and exhibits micron-sized hotspots enriched in D and ¹⁵N (Busemann et al., 2006; Nakamura-57 Messenger et al., 2006; Remusat et al., 2009). These hotspots have been interpreted as 58

remnants of interstellar organic compounds (Busemann et al., 2006; Nakamura-Messenger et al., 2006) or evidence of an early irradiation of the solar system (Remusat et al., 2009). Of note, at least for Murchison (CM chondrite), these hotspots disappear when the IOM is submitted to thermal treatment at 600°C under He atmosphere (Remusat et al., 2009).

63 Despite the abundance of available samples, much less is known about organic matter in ordinary chondrites (OCs). In contrast to most CCs, which experienced aqueous alteration 64 65 on parent bodies, most OCs experienced thermal metamorphism that may have led to the complete degradation of organics (Yang and Epstein, 1983; Alexander et al., 2010). Although 66 67 the two falls Semarkona (LL3.0) and Bishunpur (LL3.15) exhibit some signs of secondary 68 hydrous alteration (i.e. processes on the parent body, Alexander et al., 1989; Sears et al., 69 1995). Raman spectroscopy investigations on IOM have shown that Semarkona is, by far, the 70 most pristine unequilibrated ordinary chondrite (UOC). IOM experienced a metamorphic 71 temperature peak well below 250°C (Quirico et al., 2003; Busemann et al., 2007); this temperature is consistent with estimates from mineral assemblage (T<260°C - Alexander et 72 73 al., 1989) and sulfide thermometry (230°C - Zanda et al. 1995). The IOM of OCs is generally 74 richer in D than that of CCs and exhibits anticorrelated D/H and H/C ratios, in contrast to CC IOM (Alexander et al., 2007, 2010). No D- or ¹⁵N-rich hotspot has ever been reported in OC 75 76 IOM, as recently illustrated by NanoSIMS images of the IOM of the Krymka meteorite 77 (LL3.2) showing no heterogeneities in D/H (Alexander et al., 2010).

The present study reports NanoSIMS H- and N- isotopic imaging of IOM of three UOCs that experienced different thermal metamorphism: Semarkona (LL3.0), Bishunpur (LL3.15) and Grosvenor Mountains (GRO) 95502 (L3.2). In addition, we document the molecular signature and structural properties of the IOM of GRO 95502 before and after flash heating experiments. The results highlight the thermal recalcitrance of the D-rich component

- 83 of OC IOM and suggest that the original precursors of CC and OC organic matter were
- 84 different.

2. EXPERIMENTAL

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88 **2.1 Samples**

89 IOM has been isolated from three UOCs of increasing metamorphic grades: 90 Semarkona (LL3.0), Bishunpur (LL3.15), and GRO 95502 (L3.2). Semarkona and Bishunpur 91 IOM was prepared by Yang and Epstein (1983) by classical HF/HCl dissolution. This IOM 92 has been stored since then at room temperature, in a glass vial sealed by a Teflon cap and 93 protected from the sunlight. Fifty milligrams of GRO 95502 IOM (hereafter called GRO 94 IOM) were recently isolated from 20g of meteorite using a similar protocol (Piani et al., 95 2012a). An aliquot of GRO IOM was subjected to flash pyrolysis at 600°C under a He 96 atmosphere. The (organic) residue was recovered and is hereafter called GROpyr. Orgueil 97 IOM, prepared by Remusat et al. (2005), was used as a reference material.

Aliquots of these IOM samples were pressed into cleaned indium foil and gold coated (20 nm thick) to improve charge compensation during NanoSIMS imaging. Portions of GRO IOM and GROpyr samples were also finely powdered in an agate mortar and deposited on Si₃N₄ windows for Raman and XANES (X-ray absorption near edge structure) investigations.

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103 2.2 NanoSIMS settings

Isotopic images were acquired using the Cameca NanoSIMS 50 installed at the National Museum of Natural History in Paris, France. A 16 keV primary Cs⁺ beam was used to collect the secondary ions H⁻ and D⁻ during a first run (to obtain δ D images), and ¹⁶O⁻, ¹²C₂⁻ , ¹²C¹⁴N⁻, ¹²C¹⁵N⁻ and ³²S⁻ during a second run (for N/C and δ ¹⁵N images). The use of the ion ratio ¹²C¹⁴N⁻/¹²C₂⁻ reduces topographic effects on the N/C measurements (Thomen et al., 2014; Alleon et al., 2015). The primary beam was set to 8 pA for H isotope and to 3 pA for N isotope measurements, leading to spatial resolutions of about 300 and 150 nm, respectively.

111 We collected 256×256 pixel images covering $20 \times 20 \ \mu\text{m}^2$ with a raster speed of 2ms/pix. Prior 112 to each analysis, a $25 \times 25 \ \mu\text{m}^2$ presputtering was applied using a 600 pA primary current for 6 minutes (corresponding to a Cs^+ fluence of 2.2×10¹⁷ at.cm⁻²) in order to remove the gold coat. 113 114 clean the surface, and reach the sputtering steady state (Thomen et al., 2014). We used 115 Hamamatsu discrete dynode electron multipliers with a dead time of 44 ns in multicollection 116 mode. For H isotopes, the mass spectrometer was set to a mass resolving power of 4000, and to 8000 for N isotopes to resolve isobaric interferences, such as ${}^{12}C^{14}N^{-}$ from ${}^{12}C_2H_2^{-}$ and ${}^{32}S^{-}$ 117 from ${}^{16}\text{O}_2^-$. The vacuum in the analysis chamber never exceeded 5.10¹⁰ torr. 118

119 The collected NanoSIMS data were then processed with the L'Image software 120 developed by Larry Nittler, Carnegie Institution in Washington DC, USA. Each image being a 121 stack of several frames, the first step consists of aligning each frame using a correlation 122 algorithm and applying the same shift in X and Y to all the pixels of a single frame. Then, 123 ratio images can be generated. Each ratio is corrected using a calibration line (see 124 supplementary online material), determined by measuring four known reference samples 125 (which in the meantime allow checking the instrument stability): a terrestrial Type 3 kerogen, 126 a charcoal and the IOM of Orgueil and GRO 95502 (previously measured by gas source mass spectrometry by Alexander et al., (2007). Each calibration line is calculated using a linear 127 128 regression and the R program was used to determine uncertainties associated with the 129 calibration (Table S1). All the uncertainties reported in this study are one standard deviation; 130 we propagated (using quadratic sum) counting statistics uncertainties on each regions of 131 interest (ROI) and uncertainties arising from the calibration lines, reflecting the external 132 precision of our measurements (e.g., the standard deviation for independent measurements 133 repeated several times on the same sample).

134 Isotopic ratios are expressed in delta units, following the relation: $\delta(\%) = (R_{sple}/R_{std} - 1)$ 135 ×1000, with R_{sple} being the sample isotopic ratio and R_{std} a reference ratio, such as Standard

Mean Ocean Water (SMOW: $D/H= 155.76 \times 10^{-6}$) for H isotopes and air for N isotopes 136 $(^{15}N/^{14}N = 3.67 \times 10^{-3})$. Isotopic anomalies in D (or in ^{15}N) are defined as ROIs consisting of at 137 least 50 pixels (0.3 μ m²) meeting individually the requirement $|(D/H)_{pixel}-(D/H)_{average}|/\sigma > 2$ 138 (or $|({}^{15}N/{}^{14}N)_{pixel} - ({}^{15}N/{}^{14}N)_{average}|/\sigma > 2$), where (D/H)_{pixel} (or $({}^{15}N/{}^{14}N)_{pixel}$) is the isotopic ratio 139 of the pixel, (D/H)_{average} (or $({}^{15}N/{}^{14}N)_{average}$) the average ratio of the image and σ the statistical 140 141 error for each pixel (Table S2). Anomalies can be significantly positive or negative, reflecting enrichment or depletion in D (or ¹⁵N) compared to the average of the image, respectively. The 142 143 heterogeneity in the D/H ratio can be investigated using pixel histograms showing the distributions of δD (or $\delta^{15}N$) values of individual pixels within each image. The occurrence of 144 positive micron-scale anomalies can be noticed on these histograms. They result in tails on 145 146 the right side of histograms but do not affect the full width at half maximum (FWHM). Of 147 note, some of these anomalies may correspond to presolar grains (e.g. Floss and Stadermann, 148 2009). Data are reported in the supplementary online material (tables S2 and S3).

149

150 **2.3 Raman and STXM/XANES characterization**

151 Raman data were collected on powdered samples using the Renishaw INVIA 152 microspectrometer operating at IMPMC following the procedure described in (Bernard et al., 153 2008). Twenty spectra were collected on each IOM to capture their homogeneous or heterogeneous nature. Spectra were measured from 500 to 3500 cm⁻¹ at constant room 154 155 temperature using the 514.5 nm wavelength of a 50 mW Modulaser Argon laser (green laser) 156 focused on the sample through a Leica DMLM microscope with a long working distance 50X 157 objective (NA = 0.55). This configuration yields a planar resolution of $\sim 1 \mu m$ for a laser 158 power delivered at the sample surface set at around 250 µW to prevent irreversible thermal 159 damage due to laser-induced heating (Beyssac et al., 2003). A grating with 1800 lines/mm 160 disperses light and the signal was analyzed with a RENCAM CCD detector. Measurements

are performed with a circularly polarized laser using a ¹/₄ wavelength plate placed before the
microscope in order to limit polarization effects.

163 XANES data were collected using the STXM (scanning transmission X-ray 164 microscope) located on beamline 5.3.2.2 (STXM Polymer beamline - (Kilcoyne et al., 2003)) 165 at the Advanced Light Source (ALS). This beamline uses soft X-rays (250 - 600 eV) 166 generated via a bending magnet while the electron current in the storage ring is held constant 167 in top-off mode at 500 mA at a storage ring energy of 1.9 GeV. The microscope chamber is 168 evacuated to 100 mTorr after sample insertion and back-filled with He. Energy calibration is 169 accomplished using the well-resolved 3p Rydberg peak at 294.96 eV of gaseous CO₂ for the 170 C K-edge. Alignment of images of stacks and extraction of XANES spectra were done using 171 the aXis2000 software (ver2.1n) while normalization to the carbon quantity was done using 172 the Athena software package (Ravel and Newville, 2005). The C-XANES spectra shown in 173 the present study are for homogeneous organic-rich areas of several hundreds of nanometers. 174 Although radiation damage per unit of analytical information was shown to be typically 100-175 1000 times lower in STXM-based XANES spectroscopy than in TEM-based EELS 176 (Hitchcock et al., 2008), the C-XANES data shown here were collected following the 177 procedures for X-ray microscopy studies of radiation sensitive samples recommended by 178 Wang et al. (2009). Extensive databases of reference XANES spectra (measured on hundreds 179 of C-containing compounds at the C K-edge, sometimes supported by theoretical calculations 180 using multiple scattering approaches) are available (Solomon et al., 2009).

181

3. RESULTS

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185 **3.1 NanoSIMS imaging of H isotopes**

186 The spatial distributions of hydrogen isotopes in the OC IOM are remarkably different 187 from that of the Orgueil IOM (Remusat et al., 2009). Semarkona and Bishunpur IOM exhibit 188 different isotopic compositions depending on the measured area (Figure 1). Despite their D-189 rich nature, the IOM of OCs exhibits fewer D anomalies than does Orgueil IOM (Busemann 190 et al., 2006; Nakamura-Messenger et al., 2006; Remusat et al., 2009). In addition, the 191 intensities and sizes of isotopic anomalies in OC IOM differ from those in Orgueil and other 192 CC IOM (Figure 2 and table S2). Orgueil IOM exhibits numerous micron-scale D anomalies (25 over 800 μ m², with 2000‰ < δ D < 4000‰) while the average δ D value is 193 194 indistinguishable from one region to another. In contrast, only two micron-scale D anomalies 195 are detected in Semarkona IOM (one positive with $\delta D=5000\%$ and one negative with 196 $\delta D=300\%$ while entire regions may display significant D-enrichments ($\delta D=5000\%$ for 197 instance over the area 2 in Figure 1) compared to adjacent ones ($\delta D=2000$ % over the area 1 198 in Figure 1). Bishunpur IOM displays large regions with δD values ranging from 800% to 199 2700‰ and exhibits five positive (up to $\delta D=4000\%$) and three negative (around $\delta D=200\%$) micron-scale anomalies over 1200 μ m². Despite three positive (5500‰ < δ D < 6300‰) and 200 four negative (1200% $< \delta D < 1500$ %) anomalies, the GRO IOM exhibits a quite 201 homogeneous D/H ratio (considering about 600 µm² analyzed in this study), consistent with 202 203 previous measurements (Piani et al., 2012a; Piani et al., 2015). The flash pyrolysis at 600°C of this IOM induces a D-enrichment by a factor of 2.5 ($\delta D = 9000\%$) and seems to 204 205 homogenize the D/H ratio: no significant isotopic anomalous regions can be distinguished within GROpyr over 2170 μ m² surface area analyzed (Figure 3, figure S3 and Table S2). 206

207 The histograms of the δD values of individual pixels in the IOM of the OCs are 208 different from the histograms usually observed for types 1 and 2 CCs (Figure 4). For Orgueil, 209 the δD histogram is rather narrow, with asymmetry arising from the occurrence of numerous 210 D-rich hotspots on the right side (high values). The tight width of Orgueil IOM histogram 211 indicates that most of the IOM has a homogeneous isotopic ratio representing the bulk value 212 of this IOM, the hot spots only representing some material in the tail of the histogram for the 213 larger values. The contribution of the D-rich hotspot pixels to the total images of Orgueil IOM 214 has previously been shown to be small (Remusat et al., 2009). For the Semarkona and 215 Bishunpur IOM (Figure 4), the δD histograms are broader and appear bimodal (see area 2 in 216 both IOM in Figure 4). These histograms are fully consistent with the large differences in the 217 average image δD of Semarkona and Bishunpur IOM (Figure 2) and constitute suitable tools 218 to assess micron-scale isotopic distributions in IOM. Interestingly, the D-poor component in 219 Bishunpur area 2 has the same isotopic signature as the material in Bishunpur area 1. In 220 contrast, the δD histogram of the GRO IOM (Figure 4) indicates a homogeneous distribution 221 of D and the occurrence of a single main D-rich component.

222



Figure 1: NanoSIMS &D images of the IOM of Orgueil, GRO 95502, Semarkona and Bishunpur. All the images have the same size; the color scale represents corrected δD values. To illustrate the large-scale variations observed in Semarkona and Bishunpur IOM, two different areas are displayed.





Figure 2: δD values determined by NanoSIMS imaging, see also table S2. See text for ROI definition. The size of the disks represents the size of each ROI. The largest circles represent the average image values. In Bishunpur IOM, some regions identified as anomalies in one image may just correspond to "normal" area in other images.



240 241

242 Figure 3: Comparative evolution of δD NanoSIMS images for Murchison and GRO 95502 243 IOM upon 600°C flash heating. The data for Murchison were taken from Remusat et al. 244 (2009). In order to visualize the isotopic evolution, the color scale is the same before and after 245 heating. The reader is advised that we have reported here, for the unheated GRO IOM on the 246 left hand side, an image of the same areas as in figure 1, but the scale was modified to match 247 the scale of the image of GRO IOM after heating (i.e. GROpyr sample), on the right hand 248 side. For Murchison, the isotopic anomalies are destroyed and some D is lost; it results in a Ddepletion. For GRO 95502, we observe an overall D-enrichment. 249



Figure 4: pixel (frequency) histograms (pixel value distribution) of the images reported in
Figure 1. The vertical axis is in arbitrary units and represents the number of pixels having a
given δD. δD values are corrected. Bishunpur and Semarkona IOM show clearly distinct
histograms.

3.2 NanoSIMS imaging of N-isotopes

Semarkona, Bishunpur and GRO 95502 IOM do not display large-scale $\delta^{15}N$ 257 heterogeneities: each investigated IOM exhibits identical mean δ^{15} N values (within error bars) 258 over different regions (Figure 5, Figure 6 and Table S3). The Bishunpur IOM does not exhibit 259 any ¹⁵N anomalies, the Semarkona IOM only one ($\delta^{15}N$ > 200‰) and GRO IOM only two 260 (δ^{15} N> 225‰). The ¹⁵N anomalies of the Semarkona and GRO 95502 IOM are spatially 261 correlated to D anomalies (compare Figures 1 and 5). Some of these ¹⁵N anomalies exhibit 262 263 significantly lower N/C values than the bulk IOM values. In contrast the Orgueil IOM exhibits numerous micron-scale ¹⁵N positive (120‰ < δ^{15} N < 230‰) and negative (-200‰ < 264 δ^{15} N < -70‰) anomalies; they exhibit the same N/C ratios as the bulk IOM. The occurrence 265 of these anomalies in Orgueil IOM is consistent with data reported for different CCs 266 267 (Busemann et al., 2006; Nakamura-Messenger et al., 2006; Floss and Stadermann, 2009).

268 While the Bishunpur IOM shows δ^{15} N histograms similar to that of Orgueil (i.e., a 269 Gaussian curve with a FWHM of 180‰), the Semarkona and GRO 95502 IOM exhibit 270 broader histograms with FWHM of 260‰ (Figure 7). This may result from heterogeneities in 271 δ^{15} N values or, more likely, from a total N abundance that is three times lower for the IOM of 272 these chondrites compared to Bishunpur and Orgueil IOM. Such a low abundance may indeed 273 increase the measurement uncertainty of each pixel and thus artificially increase the δ^{15} N 274 variability.



Figure 5: NanoSIMS δ^{15} N images of the IOM of Orgueil, GRO 95502, Semarkona and Bishunpur. All the images have the same size; the color scale represents corrected δ^{15} N values. Theses images are located at the same places as the corresponding images in Figure 1.



Figure 6: $\delta^{15}N$ of isotopic anomalies detected in the IOM studied. The size of the disks represents the size of each ROI. The largest circles represent the average image values. Positive anomalies in Semarkona and GRO 95502 are rare and plot in the higher range of Orgueil IOM anomalies.



Figure 7: pixel (frequency) histograms (pixel value distribution) of the images reported in Figure 5. The vertical axis is in arbitrary units and represents the number of pixels having a certain δ^{15} N. Corrected δ^{15} N are reported.

290 **3.3** Evolution of the molecular structure of GRO IOM upon heating

291 The molecular evolution of OC IOM was investigated by characterizing GRO IOM 292 and GROpyr using both Raman and STXM-based XANES. The Raman signature of the GRO 293 IOM (Figure 8) is consistent with a moderately heated IOM, comparable to Krymka and 294 Inman OM (Quirico et al., 2003), with two main bands of similar intensities: a quite broad D1 band at 1340 cm⁻¹ and a composite G+D2 band centered at 1600 cm⁻¹. In contrast to the GRO 295 296 IOM, GROpyr appears structurally heterogeneous as indicated by the two different Raman 297 spectra that are shown in Figure 8. Of note, all GROpyr Raman spectra exhibit a well-298 identified G band that dominates the signal, as well as D1 and D2 bands that vary in intensity 299 depending on the degree of carbon organization. Overall, these Raman spectra point to 300 thermally evolved organics having experienced partial graphitization (Bernard et al., 2010).

301 The XANES spectrum of GRO IOM exhibits two prominent absorption peaks at 285.3 302 and 288.6 eV corresponding to electronic transitions of aromatic and/or olefinic carbon (C=C) 303 and of carbon in carboxylic functional groups (COOH), respectively. The presence of 304 aliphatic carbon (CH_n) can also be seen at 287.5 eV. Despite the absence of ketonic carbon 305 (peak at 286.5 eV), this spectrum appears quite similar to that of CC IOM such as Orgueil or 306 Murchison IOM (De Gregorio et al., 2013; Le Guillou et al., 2014). GROpyr contains 307 significantly more aromatic/olefinic carbons than the GRO IOM together with a lower 308 concentration of aliphatic and carboxylic carbons. The evolution of the XANES signal from 309 GRO IOM to GROpyr is consistent with Raman data and shows molecular evolution during 310 partial graphitization (Bernard et al., 2010).

Taken together, Raman and XANES signatures highlight the similar molecular and structural evolution undergone by both CC IOM (Bonal et al., 2007; Busemann et al., 2007; Cody et al., 2005; Remusat et al., 2008) and OC IOM upon heating (flash pyrolysis at 600°C).



Figure 8: Structural and molecular signature evolution of GRO IOM under thermal stress. A: Raman spectra show that GRO IOM is becoming more organized, although it gets heterogeneous. B: XANES spectra of GRO IOM and GROpyr. XANES spectra show that upon heating, GRO IOM is becoming more aromatic and is losing aliphatic C. In addition, COOH groups are less abundant.

4. DISCUSSION

- 324
- 325 **4.1 D and ¹⁵N distributions in OC IOM**

The most D-rich organic moieties in the Orgueil IOM are organic radicals (Delpoux et al., 2010), which are heterogeneously distributed in the IOM (Binet et al., 2004). The CI, CM and CR IOM have been described as assemblage of moderately D-rich organic matter associated with organic radicals extremely rich in D that constitute less than 20 vol % of the total IOM (Remusat et al., 2009). The clusters rich in organic radicals then appear as micronsize D-rich hotspots in NanoSIMS images (Remusat et al., 2009).

332 In contrast, very few micron-scale D anomalies have been observed in the OC IOM 333 investigated in the present study, consistent with observations reported for the Krymka (LL 334 3.2) IOM (Alexander et al., 2010). Nevertheless, the D distribution in OC IOM is not 335 homogeneous. Semarkona and Bishunpur IOM seem to be constituted by at least two 336 components that can be resolved at the scale of several microns (Figure 1 and 4). These 337 organic components may have experienced different conditions on the parent body 338 (Alexander et al. 2010) or may have originally been different, i.e. they may have had distinct 339 δD signatures before being accreted. It has been shown that large variations of clay mineral 340 isotope signatures and heterogeneities of organic isotopic signatures are not correlated within 341 the matrix of Semarkona (Piani et al., 2015). This observation strengthens the scenario of the 342 accretion of distinct organic precursors having different isotopic signatures.

The occurrence of large domains with distinct D/H is intriguing. Analytical bias during NanoSIMS imaging can be ruled out as such distribution is not observed in terrestrial standards or Orgueil IOM measured under the same experimental conditions. In addition, the average values obtained for Semarkona and Bishunpur IOM are consistent with previous measurements (Alexander et al. 2007), thereby disproving the possibility of an HF/HCl dissolution artifact (it is unlikely that HF/HCl treatments would induce such D-transfer between organic moieties). The occurrence of these large domains with distinct D/H may result from the intrinsic nature of the different components of the IOM. OCs exhibit micrometric organic grains together with a diffuse organic component within the matrix, as observed in CCs (Le Guillou et al., 2014; Piani et al., 2015). During HF/HCl dissolution, organic components with similar affinities may gather together forming 10-20 micron large aggregates that may escape destruction during the centrifugation and drying steps.

355 Despite variable high D/H values, the OC IOM investigated present an almost homogeneous N-isotope composition close to $\delta^{15}N=0$ %. Alexander et al. (2007; 2010) 356 357 reported very large D-enrichment in bulk IOM from various OCs, associated with Earth-like 358 N isotopic compositions using conventional isotope ratio mass spectrometry. This is in 359 contrast to the CCs where the D-richest IOM also exhibit large ¹⁵N-enrichments. Again, such 360 differences may not be explained by different parent body evolution but rather by the accretion of both D- and ¹⁵N-rich organics in CRs and a few CMs while only D-rich organics 361 were accreted in OCs. The very few ¹⁵N anomalies in OC IOM exhibit large D-enrichments 362 (making them both D- and ¹⁵N-rich hotspots). Ion-molecule or grain surface chemistry in cold 363 environments could explain both D- and ¹⁵N-enrichments, their scarcity possibly resulting 364 365 from their sensitivity to the thermal metamorphism that occurred on OC parent bodies.

366

367 **4.2 Thermal evolution of OC IOM**

368 While the bulk δ^{15} N signatures do not appear to be sensitive to temperature conditions, 369 the present investigations of three UOCs of increasing metamorphic grades (Semarkona 370 (LL3.0), Bishunpur (LL3.15) and GRO 95502 (L3.2)) demonstrate an increase of δ D values 371 with increasing metamorphism. In contrast, the D/H and H/C ratios of CC IOM decrease with 372 increasing metamorphism (Alexander et al., 2007). In other words, OC and CC IOM exhibit opposite trends in the D/H vs. H/C plot (Alexander et al., 2010), even though the structural
evolution of OC and CC IOM are similar with increasing temperature (Quirico et al., 2003;
Bonal et al., 2007).

376 Thermal degradation experiments (flash heating) performed on Murchison IOM 377 demonstrated that D-rich hotspots do not sustain temperature (Remusat et al., 2009). Such 378 result makes sense if hotspots concentrate D-rich organic radicals that can be easily thermally 379 destabilized (Remusat et al., 2009). Furthermore, the IOM of the Kainsaz chondrite (CO3.6), 380 which lacks D-rich hotspots, contains a very low amount of radicals (Remusat et al., 2008; 381 Remusat et al., 2009). In contrast, the present study demonstrates that the D/H ratio of GRO 382 IOM increases with increasing temperature (GROpyr D/H is twice as large as GRO IOM) 383 despite a structural and molecular evolution similar to that of CC IOM. Of the two organic 384 components described in 4.1, one is likely thermally recalcitrant (the D-rich one) while the 385 second evolves into graphitized material through H loss (its H/C decreases with increasing 386 temperature). Indeed, as indicated by Raman and STXM-based XANES spectroscopy, the 387 GRO IOM undergoes a structural reorganization concomitant with an increase of its 388 aromatic/aliphatic carbon ratio as temperature increases. This evolution, typical of the 389 carbonization process (Bernard et al., 2010), is also observed for CC IOM affected by thermal 390 metamorphism (Quirico et al., 2003; Bonal et al., 2007; Busemann et al., 2007, Remusat et 391 al., 2008).

The present contribution thus demonstrates that the organic D-rich reservoirs in OC and CC IOM are different. While CC IOM loses its most D-rich organic reservoir upon heating, OC IOM loses its D-poor organic H reservoir. This is consistent with the evolution from a bimodal D/H histogram, as seen in the Semarkona and Bishunpur IOM, to a more homogeneous distribution in the IOM of the more heated GRO 95502 (Figure 4). This is also

consistent with NanoSIMS data reported for the IOM of the Krymka 3.2 OC (Alexander et al.,2010).

In summary, in contrast to that of CC IOM, the D-rich component of OC IOM is not borne by organic radicals but rather by thermally recalcitrant organic moieties. This recalcitrance would explain the evolution reported by Alexander et al. (2007; 2010) for bulk OC IOM in the D/H vs. H/C plot, i.e. the increase of the D/H ratio of OC IOM may directly result from an increase of metamorphic temperature experienced by OC parent bodies.

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405 **4.3 Origin of the D-rich organic reservoir in OCs**

406 Alexander et al. (2010) suggested that the D-rich component of UOC IOM results 407 from isotopic exchange between IOM and D-rich water, such as the water reported in UOCs 408 by Deloule and Robert (1995). This water would have been enriched in D through distillation 409 process resulting from the progressive reduction of water by reaction with Fe. Yet, Piani et al. 410 (2015) recently reported the absence of spatial correlation between IOM and water D/H ratios 411 as well as huge heterogeneities in the δD values of Semarkona phyllosilicates (up to δD = 412 10,000 ‰) that may not result from distillation processes. Aléon (2010) suggested an 413 alternative hypothesis: he interpreted the D-enrichment of OC IOM as resulting from the 414 presence of presolar organics synthesized in the interstellar medium (ISM). In this model, the 415 D/H ratio of OC IOM results from mixing between IOM similar to that of Murchison and extremely D-rich ISM organics (1 at.% at most), with no significant ¹⁵N-enrichment relative 416 417 to solar system organic matter. Hence, according to Aléon (2010) and Piani et al. (2015), both 418 D-rich organics and D-rich water ice from primitive reservoir(s) would have been accreted 419 and preserved within OC parent-bodies.

In Figure 9, the data collected in the present study are compared to previousmeasurements of CC and OC IOM, IDPs and Hale-Bopp HCN as well as models showing

fractionation due to ion-molecule reactions (IMR-fl) and mixing between a component akin to CM chondrites IOM (Murchison) and observed molecules in the interstellar medium (Aléon, 2010 and references therein). H and N isotopic compositions are expressed as isotopic fractionation factors (α H and α N) between organic matter and the protosolar gas, the composition of which is D/H = 21×10⁻⁶ (Geiss and Reeves, 1981) and ¹⁵N/¹⁴N = 2.36×10⁻³ (Meibom et al., 2007). The Genesis mission since reported that protosolar ¹⁵N/¹⁴N = 2.27(± 0.03) ×10⁻³ (Marty et al., 2011).

429 The present data are consistent with previous OC IOM data and fall on the mixing 430 curve proposed by Aléon (2010). Most of the D-enrichments reported here are not associated with any significant enrichment in ¹⁵N. At the micro-scale, no D anomalies correspond to the 431 432 ISM organic component with D/H=0.1 predicted by Aléon (2010). Thus, if mixing has occurred, it has to be at a very fine scale, below a few tens of nanometers, i.e. at the molecular 433 434 scale. Hence, OC IOM contains interstellar molecular moieties or was formed from ISM 435 precursors that were incorporated into the OC IOM during solar system formation. Of note, D and ¹⁵N anomalies of OC IOM follow the trend defined by the simultaneous H and N isotopic 436 437 fractionation by ion/molecule reactions occurring in the protosolar nebula or in the parent 438 molecular cloud (IMR-fl, Figure 9); hence, they may have formed in the solar system nebula 439 by ion/molecule reactions (Aléon, 2010).



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Figure 9: Comparison of the isotopic composition of OC IOM (this study) with the model by 444 445 Aléon (2010). The N isotopic fractionation between organics and the protosolar gas is 446 reported versus the H isotopic fractionation. Black dots represent sample of group 1 as 447 defined by Aléon: IOM in CC and their hotspots, values taken from the literature until 2010. 448 In open black dots are reported bulk IOM of unequilibrated OC (Alexander et al. 2007). 449 Curves describe mixing between observed interstellar molecules (Observed ISM) and typical 450 CM chondrite IOM with: isochemical mixing (plain curve), mixing with Murchison CHON 451 and ISM HCN (dashed curves) and with Murchison HCN and ISM CHON (doted curved). 452 ISM values are taken from Millar et al. (1989), Turner (2001) and Roueff et al. (2007). 453 Details and references can be found in Aléon (2010). Our data for OC IOM are reported as 454 red open diamonds. It must be noted that three hotspots observed in Semarkona and GRO 455 95502 fit with the ion/molecule reaction fractionation line (IMR-fl) defined by (Aléon, 2010). 456

5. CONCLUSION

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The present study shows that the D and ¹⁵N distributions in CC and OC IOM are 459 460 different. The OC IOM does not exhibit numerous micron-scale isotopic anomalies (both D-461 and ¹⁵N-rich), even in the least metamorphosed, Semarkona. In addition, OC IOM appears to 462 be composed of two main isotopic reservoirs, the D-richer being thermally recalcitrant. 463 Thermal degradation experiments (flash heating) indeed demonstrate that the D/H ratio of OC 464 IOM increases with increasing temperature despite a structural and molecular evolution 465 similar to that of CC IOM as indicated by Raman and STXM-based XANES spectroscopy. 466 Based on these observations, we suggest that CCs and OCs did not accrete the same organic 467 precursors, as a consequence of spatial and/or temporal heterogeneities in the solar nebula. 468 This indicates that the organic content of the protosolar nebula was probably evolving during 469 parent body formation. The common organic precursor concept should be revised, as CC, OC, 470 and enstatite chondrites appear to contain IOM with different properties (Alexander et al., 471 2007; Piani et al., 2012b). We suggest that the D-rich, thermally recalcitrant component of 472 OC IOM may have formed in the ISM, thus supporting the vision of Aléon (2010) stating that 473 OC IOM is "polluted" by minute amount of ISM organics.

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