

Early stages of core segregation recorded by Fe isotopes in an asteroidal mantle

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Abstract :

Ureilite meteorites are achondrites that are debris of the mantle of a now disrupted differentiated asteroid rich in carbon. They provide a unique opportunity to study the differentiation processes of such a body. We analyzed the iron isotopic compositions of 30 samples from the Ureilite Parent Body (UPB) including 29 unbrecciated ureilites and one ureilitic trachyandesite (ALM-A) which is at present the sole large crustal sample of the UPB. The $\delta^{56}\text{Fe}$ of the whole rocks fall within a restricted range, from 0.01 to 0.11‰, with an average of $+0.056 \pm 0.008\text{‰} + 0.056 \pm 0.008\text{‰}$, which is significantly higher than that of chondrites. We show that this difference can be ascribed to the segregation of S-rich metallic melts at low degrees of melting at a temperature close to the Fe–FeS eutectic, and certainly before the onset of the melting of the silicates (View the MathML source <1100°C), in agreement with the marked S depletions, and the siderophile element abundances of the ureilites. These results point to an efficient segregation of S-rich metallic melts during the differentiation of small terrestrial bodies.

Highlights

► Ureilites displays $\delta^{56}\text{Fe}$ values higher than average chondrite. ► Segregation of Fe-sulfide melts explains the high $\delta^{56}\text{Fe}$ values in ureilites. ► Formation of a core can begin at very low degrees of melting through the circulation of a Fe–S melt through a silicate mantle.

Keywords : iron isotopes, achondrite, ureilite, core

1. Introduction

51 One of the most significant issues for understanding the differentiation of telluric bodies is
52 when and how their cores formed. Trace element abundances and W isotopic compositions obtained
53 on a series of iron meteorites indicate that the cores in the first differentiated planetesimals aggregated
54 metallic melts with various S contents, and formed from ca. 0.6 to 3 Myrs after the first solids
55 condensed in the Solar System (e.g., Blichert-Toft et al., 2010; Kruijer et al., 2014 and references
56 therein). The mechanisms of core formation are still highly debated, especially for the early stages of
57 the process. The percolation of Fe-S melts into an olivine matrix has been suggested and
58 experimentally investigated, but the contribution of such melts to the cores of the planetary embryos or
59 of larger bodies, is a matter of debate (Bruhn et al., 2000; Yoshino et al., 2003; Bagdassarov et al.,
60 2009; Rushmer and Petford, 2011; Watson and Roberts, 2011). However, important constraints on the

61 early stages of the differentiation of some small bodies are potentially recorded in their mantles, which
62 can be investigated using meteorite samples.

63 Ureilites are remnants of the mantle of a now-disrupted carbon-rich body (Downes et al.,
64 2008), and are among the most common achondrites in the meteorite collections (about 400 ureilites
65 are currently reported in the Meteoritical Bulletin Database). They are ultramafic achondrites (e.g.,
66 Mittlefehldt et al., 1998) which are widely considered as mantle restites (Scott et al., 1993) formed
67 after extraction of magmas and S-rich metallic melts. The vast majority of them are unbrecciated.
68 They are coarse-grained peridotites consisting chiefly of olivine and pyroxenes (pigeonite, and more
69 rarely augite and orthopyroxene), abundant carbon (< 7 wt%, graphite and diamond), with accessory
70 metal and sulfides. Olivine grains classically have distinctive iron-depleted rims and veins (Fig. 1).
71 They were formed by local reduction reactions with adjacent carbon. These rims give important
72 constraints on the Ureilite Parent Body's (UPB) history. The Fe-Mg zonings of the olivines indicate
73 that ureilites equilibrated at high temperatures (1200-1300°C), and subsequently rapidly cooled down
74 (~2-6°C/h), consistent with impact-excavation or disruption of the UPB (e.g., Myiamoto et al., 1985).
75 Except these rims and veins in olivine, silicates (i.e., olivine cores and pyroxenes) are quite uniform in
76 Mg# number [= 100 Mg/(Mg+Fe), atomic] within any given ureilite. However, their compositions
77 among samples show a huge variation, as shown by the olivine-core Mg# (= forsterite content), which
78 range from 74 to 97. The Fe/Mn ratios (ranging from 3 to 57) and $\Delta^{17}\text{O}$ values (= -0.2 to -2.5 ‰) are
79 correlated with this parameter (e.g., Clayton and Mayeda, 1996; Mittlefehldt et al., 1998; Fig. 2). The
80 origin of these variations is a matter of discussions, but no consensus emerges at present. Certainly,
81 they cannot be explained by igneous fractionation (e.g., Mittlefehldt et al., 1998). Instead, it has been
82 often argued that the Mg# range was ascribed to redox processes during the differentiation of the UPB
83 or alternatively is inherited from the pre-igneous (nebular) history of the accreted materials (see
84 Warren (2012) and Goodrich et al. (2013a) for extensive discussions of the current models on its
85 origin).

86 A score of ureilites are polymict breccias consisting of debris of these peridotites, various
87 chondritic clasts, and some feldspathic clasts which are remnants of the UPB's lavas (e.g., Cohen et

88 al., 2004; Downes et al., 2008). A large clast (ALM-A) has been recently discovered among the
89 numerous stones of the Almahata Sitta fall, and demonstrate that at least part of these melts was
90 trachyandesitic (Bischoff et al., 2014).

91 Unlike asteroids such as Vesta whose O isotopic compositions were homogenized by a magma
92 ocean (Greenwood et al., 2005), the wide range of $\Delta^{17}\text{O}$ values displayed by the ureilites demonstrates
93 that the UPB was never totally melted (Clayton and Mayeda, 1996). Nevertheless, it was heated
94 enough to generate magmas (e.g. Cohen et al., 2004; Bischoff et al., 2014) and a small S-rich core
95 (Warren et al., 2006; Rankenburg et al., 2008; Goodrich et al., 2013b).

96 Iron isotope ratios can track the involvement of iron sulfides during the early stages of the
97 formation of the asteroidal cores. Marked enrichments in the lighter iron isotope are found in iron
98 sulfides formed at ca. 1000°C or below (Schuessler et al., 2007; Polyakov and Soultanov, 2011; Wang
99 et al., 2014), whereas no iron isotope fractionation has been detected between iron sulfide, metal and
100 silicate melts at ca. 1250°C (e.g., Hin et al., 2012). Here, we report high-precision iron isotope
101 compositions of 30 meteorites from the UPB, and show that its mantle displays iron isotopic
102 compositions heavier than average chondrite. This indicates that a significant fraction of the sulfides
103 was efficiently segregated during the early stages of the melting, possibly before the melting of the
104 silicates. Therefore, core formation in an asteroid can begin at very low degrees of partial melting
105 through the circulation of a Fe-S melt through a silicate mantle.

106

107 **2. Samples and analytical procedures**

108 We analyzed 30 meteorites from the UPB (29 unbrecciated ureilites and a unique lava sample,
109 Table 1). Samples were kindly provided by the NASA meteorite working group (MWG), the National
110 Institute of Polar Research (NIPR), the Ecole Normale Supérieure de Lyon (ENSL), the Westfälische
111 Wilhelms-Universität Münster, the Washington University in Saint Louis, and finally from the first
112 author's collection. The samples from Antarctica have been extensively studied by different teams.
113 The compositions of their phases have been repeatedly determined and their chemical features are well

114 known (the references are too numerous to be cited here). Our meteorites from Sahara have been less
115 studied, and for most of them, only their descriptions in the Meteoritical Bulletin are available. They
116 are all regular ureilites. We have examined polished sections of all these samples to confirm their main
117 petrographical features, and determine the compositions of their silicates. The Saharan samples were
118 selected in order to complement the range of compositions displayed by the Antarctic ones. Although
119 the terrestrial weathering is pervasive in hot desert samples, the samples we selected were not severely
120 weathered, and display partly preserved metal and sulfides.

121 The major-element compositions of the phases were determined by electron microprobe
122 analysis using a Cameca SX100 at Service Commun de Microsonde Ouest (SCMO), Plouzané, or a
123 JEOL JXA8200 at NIPR (Table 2). Minerals and metal standards were used for calibration. All
124 analyses used wavelength dispersive spectrometers at 15 kV accelerating voltage, 10–30 nA beam
125 current at NIPR, and 20 kV accelerating voltage, 40 nA beam current with the same counting time as
126 Pierre Hudon and David Mittlefehdt (in Downes et al., 2008) at Plouzané .

127 Fragments (each of 300-1,000 mg) were crushed to a homogeneous fine powder using a boron
128 carbide mortar and pestle. Twenty mg of crushed material was digested by sequential mixtures of
129 HF/HNO₃, HNO₃ and HCl for the samples analyzed in Brest and in HF/HNO₃ and HCl for the samples
130 analyzed in St Louis. These procedures allow a perfect dissolution of all the phases except graphite
131 and diamond, which are Fe-free, have no effect for our study.

132 Aliquots of the solutions were used for the determination of the Fe and Mg concentrations of
133 in the solutions were determined by ICP-AES (inductively coupled plasma-atomic emission
134 spectrometry) using a Horiba Jobin Yvon Ultima 2 spectrometer in Plouzané, and the Mg# numbers
135 are given in Table 1. Iron in the dissolved samples was then purified by ion-exchange chromatography
136 procedure previously described (Wang et al., 2012). The purification was repeated twice, thus
137 ensuring the removal of all the matrix elements. A majority of the Fe isotopic compositions were
138 measured on the Thermo Electron Neptune multi collector-inductively coupled plasma-mass
139 spectrometer housed at Ifremer, Plouzané, while the rest was analyzed in Saint Louis. In both Saint
140 Louis and Brest the instrument was operated in high resolution mode to resolve isobaric interferences,

141 such as ArO on ^{56}Fe , ArOH on ^{57}Fe , and ArN on ^{54}Fe . We also used the sample-standard bracketing
 142 technique to correct for instrumental mass discrimination by normalizing Fe-isotope ratios to the
 143 average measured composition of the standard that was run before and after the sample. In Plouzané,
 144 we performed an additional correction using Ni as an internal standard for mass bias correction
 145 (Dauphas and Rouxel, 2006). Fe isotope values are reported relative to the standard IRMM-14 using
 146 the δ -notation:

$$147 \quad \delta^x\text{Fe} = 1000 \left[\left(\frac{^x\text{Fe}/^{54}\text{Fe}}{\text{sample}} \right) / \left(\frac{^x\text{Fe}/^{54}\text{Fe}}{\text{IRMM-14}} \right) - 1 \right] \text{ where } x=56 \text{ or } 57.$$

148 Results for several georeference materials are similar to previously reported analyses (Table 3).
 149 Errors are reported as 2 standard errors, which are calculated as the standard deviation divided by the
 150 square root of the total number of analyses and multiplied by the Student's t-value for the relevant
 151 degree of freedom at the 95% confidence level. Results for the samples are listed in Table 4.

152

153 3. Results

154 The $\delta^{56}\text{Fe}$ values obtained for 30 meteorites from the UPB, fall within a restricted range, from
 155 0.01 to 0.11‰ (Fig. 3, Table 4). The mean of all these samples is significantly heavier ($\delta^{56}\text{Fe} = 0.056$
 156 ± 0.008 ‰, $n=30$) than that of primitive meteorites, such as carbonaceous, ordinary and enstatite
 157 chondrites ($\delta^{56}\text{Fe} = 0.005 \pm 0.008$ ‰, $n=42$, Craddock et al., 2013). The heavy Fe isotope composition
 158 displayed by the ureilites is striking because the early Solar Nebula was well homogenized for Fe
 159 isotopes (Schoenberg and von Blanckenburg, 2006; Wang et al., 2013), and because the Fe isotope
 160 compositions of the bulk silicate portions of most differentiated bodies investigated so far are
 161 chondritic [e.g., the Earth (Craddock et al., 2013), Mars and Vesta (Poitrasson et al., 2004; Weyer et
 162 al., 2005; Schoenberg and von Blanckenburg, 2006; Wang et al., 2012), and probably the Moon
 163 (Weyer et al., 2005; Halliday et al., 2013)]. One noteworthy exception is the angrite parent body
 164 whose lavas display high $\delta^{56}\text{Fe}$ ratios (≈ 0.12 ‰, Wang et al., 2012).

165

166 **4. Discussion**

167 **4.1. Iron inventory in ureilites**

168 Iron in ureilites is hosted by many phases: silicates (olivine and pyroxenes), metal, sulfides,
169 phosphides, carbide (cohenite), and their weathering products (Fig. 1). It is very difficult to obtain a
170 reliable picture of the iron budget in ureilites from modal analyses. Goodrich et al. (2014) estimated
171 that ureilites contain ~1-3 vol.% of grain boundary metal, and less sulfides, in agreement with the
172 abundances of metal and sulfur determined by chemistry (Vdovykin, 1970; Wänke et al., 1972;
173 Hintenberger et al., 1978; Yanai et al., 1995). We used the standard wet chemical analyses compiled
174 by Yanai et al. (1995) to estimate the proportions of Fe hosted by silicates, metal and troilite in
175 ureilites (Fig. 4). Our calculations indicate that silicates are the main hosts of Fe, and represent more
176 than 70 % of the budget of this element. Metal (14-29 % of the Fe budget) and sulfides (< 8 % of the
177 Fe budget) are clearly subordinate hosts. These conclusions are strengthened by the striking
178 correlation between the Mg# numbers (=100 x Mg/(Mg + Fe), atomic) of our samples and the
179 composition of their olivine cores, which are representative of the Mg# numbers of their silicates (Fig.
180 5).

181 **4.2. Origin of the high $\delta^{56}\text{Fe}$ values in ureilites**

182 The high $\delta^{56}\text{Fe}$ values in ureilites cannot be the result of terrestrial weathering. Alteration of
183 sulfides in meteorites is accompanied by the release of light Fe. Heavier Fe isotopic compositions have
184 been measured in weathered ordinary chondrite finds compared to fresh falls. However, the increase is
185 rather limited, about +0.07 ‰ of the $\delta^{56}\text{Fe}$ were observed only for the most weathered L and H
186 chondrites (Saunier et al., 2010). Ureilites are less prone to such an effect because sulfides control here
187 a much lesser proportion of Fe than in L and H chondrites (Fig. 4). Although the alteration of ureilite
188 finds is pervasive, our samples were not extensively weathered, and still displayed metal and sulfides.
189 Indeed, the fresh fall samples (Novo Urey and Almahata Sitta) display the same isotopic compositions
190 as find samples (Tables 1 and 4).

191 Because the Fe isotopic compositions are certainly pristine, they could be inherited from the
192 materials which accreted to form the UPB, or alternatively they could result of the differentiation of
193 the UPB. A heavy Fe isotopic composition could be observed in bodies that suffered evaporative loss
194 of volatile elements following high-energy impacts during accretion. This explanation was initially
195 proposed for the Earth and the Moon by Poitrasson et al. (2004) but is controversial for these bodies
196 (e.g., Halliday, 2013). It is probably valid for the angrite parent body (APB), where a strong depletion
197 in volatile elements such as the alkalis, is accompanied by heavy Fe and Si isotopic compositions
198 (Wang et al., 2012; Pringle et al., 2014). The UPB did not accrete from volatile depleted materials,
199 because the magmas generated by the melting of its mantle were alkali and volatile-rich as shown by
200 the compositions of the plagioclases in the feldspathic clasts found in polymict ureilites and the
201 geochemistry of ALM-A, a trachyandesitic lava from the UPB (e.g., Cohen et al., 2004; Bischoff et
202 al., 2014). Nevertheless, losses of volatile elements were possible during the disruption of the UPB, as
203 shown previously for Zn (Moynier et al., 2010). However, the lack of correlation between $\delta^{56}\text{Fe}$ values
204 and abundances of volatile elements (e.g., Zn, Rb, Li, not shown) rules out this possibility for Fe.

205 The high $\delta^{56}\text{Fe}$ values in ureilites cannot be created by either the extraction of silicate melts, or
206 by possible high temperature redox processes. Partial melting does not generate residues with $\delta^{56}\text{Fe}$
207 values significantly distinct than those of the initial mantle (Weyer et al., 2005; Craddock et al., 2013).
208 High $\delta^{56}\text{Fe}$ values in bulk rocks are certainly not the result of the reduction reactions that produced the
209 Fe-depleted rims of the olivine during the disruption of the body. Although, the possibility of local
210 isotopic heterogeneities in olivine have not yet been investigated, the possibility of a detectable effect
211 is unlikely: 1/ the relative volume of the Fe-depleted rims is always limited (Fig. 1), and consequently
212 the impact of these rims on the Fe budget is probably negligible; 2/ the formation of the Fe-depleted
213 olivine rims was a very fast process (e.g., Myiamoto et al., 1985), and a noticeable metal loss during
214 the post-disruption cooling of the ureilites is not plausible; 3/ ALM-A, the ureilitic trachyandesite,
215 displays a high $\delta^{56}\text{Fe}$ value which indicates that this feature predates the disruption of the body.
216 Moreover, the high $\delta^{56}\text{Fe}$ values displayed by the ureilites are certainly not generated by the processes

217 that produced the striking range of olivine-core compositions. The lack of correlation between the Fe
 218 isotope compositions of ureilites and their olivine-core compositions demonstrates that both features
 219 are unrelated (Fig. 2).

220 Alternatively, removal of S-rich metallic melts could explain the high $\delta^{56}\text{Fe}$ values in ureilites.
 221 When a chondritic assemblage is heated, melting begins with Fe-sulfide (troilite) and metal at a
 222 temperature close to 980°C defined by the eutectic in the Fe-FeS system (Keil, 2000). Since at this
 223 temperature FeS displays a marked enrichment in the lighter Fe isotope (Schuessler et al., 2007,
 224 Polyakov and Soultanov, 2011; Wang et al., 2014), an increase of the $\delta^{56}\text{Fe}$ value of the solid residue
 225 is anticipated. If the metallic melt is efficiently removed from the solid residue (fractional melting), all
 226 melting happens at the eutectic until either metal or troilite is exhausted. The effect on the isotopic
 227 compositions of the residues can be easily estimated.

228 The eutectic in the Fe-FeS system contains about 31.6 wt% S and 68.4 wt% Fe (e.g., Kullerud,
 229 1963), and is equivalent to about 85 wt% troilite +15 wt% metal. We calculate the $\Delta^{56}\text{Fe}_{\text{metallic melt-silicate}}$
 230 using these proportions, the $\Delta^{56}\text{Fe}_{\text{troilite-silicate}}$ value (-0.21 ‰) selected by Wang et al. (2014) and the
 231 $\Delta^{56}\text{Fe}_{\text{metal-troilite}}$ at ca. 1250 K (+0.1 - +0.2 ‰) estimated from theoretical calculations (Polyakov et al.,
 232 2007; Polyakov and Soultanov, 2011). For the calculations presented in Figure 6, we used a
 233 $\Delta^{56}\text{Fe}_{\text{metallic melt-silicate}} = -0.18$ ‰, and assumed that $\Delta^{56}\text{Fe}_{\text{metallic melt-silicate}} \approx \Delta^{56}\text{Fe}_{\text{metallic melt-residue}}$.

234 The composition of a source after extraction of a S-rich metallic melt at the eutectic were
 235 calculated using an isotope mass balance:

$$236 \quad \delta^{56}\text{Fe}_{\text{source}} = \delta^{56}\text{Fe}_{\text{metallic melt}} \cdot f_{\text{Fe}_{\text{metallic melt}}} + \delta^{56}\text{Fe}_{\text{residue}} \cdot f_{\text{Fe}_{\text{residue}}} \quad (1)$$

237 where f is the mass fraction of ^{54}Fe in the metallic melt or in the solid residue. This equation
 238 can be recast in terms of the residue:

$$239 \quad \delta^{56}\text{Fe}_{\text{source}} = (\Delta^{56}\text{Fe}_{\text{metallic melt-residue}} + \delta^{56}\text{Fe}_{\text{residue}}) (1 - f_{\text{Fe}_{\text{residue}}}) + \delta^{56}\text{Fe}_{\text{residue}} \cdot f_{\text{Fe}_{\text{residue}}} \quad (2)$$

240 And consequently:

241
$$\delta^{56}\text{Fe}_{\text{residue}} = \delta^{56}\text{Fe}_{\text{source}} - \Delta^{56}\text{Fe}_{\text{metallic melt-residue}} + \Delta^{56}\text{Fe}_{\text{metallic melt-residue}} \cdot f_{\text{Fe}_{\text{residue}}} \quad (3)$$

242 Calculations show that detectable shift of the Fe isotopic compositions is possible, depending
 243 on the initial Fe and S contents of the starting assemblage (Fig. 6). If the starting assemblage is Fe-rich
 244 and displays moderate S contents (e.g., like a H chondrite), segregation of the metallic melt produces
 245 at best a marginal shift of the isotopic composition of the residue. In contrast, for S-rich chondritic
 246 assemblages, an increase of ca. 0.1 ‰ of the $\delta^{56}\text{Fe}$ can be achieved by this process.

247 If the metallic melt is not removed from the residue (batch melting), the degree of melting
 248 increases with the temperature. The composition of the metallic melt is consequently variable, and the
 249 situation becomes much more complex (melting of the silicates, involvement of carbon...). More
 250 importantly, fractionation of Fe isotopes between S-rich metallic melts and silicates becomes much
 251 less detectable with increasing temperature, and is usually supposed negligible considering our present
 252 level of precision, above 1250°C as determined through experimental studies (Poitrasson et al., 2009;
 253 Hin et al., 2012). [Notice that this view has been recently challenged by Shahar et al. (2015) whose
 254 results indicate significant fractionations between metallic melts and melted silicates at 1600°C.
 255 However, the magnitude and direction of these isotopic fractionations are not consistent with the
 256 available analyses on magmatic irons (bulk samples) and the results presented here. A discussion of
 257 these results and previous ones is beyond the scope of this paper. Additional works are necessary to
 258 confirm these experimental results and to evaluate the effects of B or Sn used in previous
 259 experiments.]

260 Siderophile element systematics (e.g. platinum group elements) demonstrate that ureilites
 261 experienced separation of S-rich metallic melts, with possibly entrainment of metal (Warren et al.,
 262 2006; Rankenburg et al., 2008; Goodrich et al., 2013b), but the process (batch or fractional) and
 263 temperatures of segregation are not well constrained by these data. The heavy Fe isotopic
 264 compositions and the low S contents of the ureilites are consistent with the separation of S-rich
 265 metallic melts (Fig. 6), and indicate that the removal of these melts was efficient probably at a
 266 temperature close to the eutectic, and certainly before the onset of the melting of the silicates (< 1100

267 °C). Indeed, ALM-A, the sole ureilitic lava sample available at present (Bischoff et al., 2014), displays
 268 a high $\delta^{56}\text{Fe}$ value (Fig. 3), similar to the highest values measured on ureilites. Although partial
 269 melting can generate melts with Fe isotopic compositions heavier than their sources as exemplified by
 270 terrestrial basalts (e.g., Teng et al., 2013), this effect is negligible for mantles with low $\text{Fe}^{3+}/\text{Fe}_{\text{total}}$
 271 ratios (Dauphas et al., 2014), like ureilites (Goodrich et al., 2013c). Consequently, ALM-A strongly
 272 suggests that the UPB's mantle acquired its specific isotopic composition early, before it was heated
 273 enough to generate magmas.

274 **4.3. Comparison with mantles of other small bodies.**

275 The heavy Fe isotopic compositions displayed by ureilites contrast with those of the
 276 brachinites and the olivines from the Main Group Pallasites, the samples of the other asteroidal
 277 mantles for which data are available (Fig. 3). Brachinites were possibly derived from multiple parent
 278 bodies. They are mantle restites similar to ureilites, but are more ferroan than the latter (Keil, 2014).
 279 GRA 06128/29, a brachinitic melt shows a light Fe isotopic composition ($\delta^{56}\text{Fe} = -0.08 \pm 0.06 \text{ ‰}$),
 280 which was ascribed to the involvement of Fe-sulfides during magma genesis (Wang et al., 2014).
 281 Probably because of the high Fe content of their protoliths, the average composition of the brachinites
 282 does not deviate from the chondritic value ($\delta^{56}\text{Fe} = 0.01 \pm 0.02 \text{ ‰}$, $n=7$, Wang et al., 2014), although
 283 S-rich metallic melts and sulfides were certainly involved during their differentiation (Day et al.,
 284 2012). Main Group Pallasites (MGPs) are mixtures of mantle-derived olivines and core-derived metal
 285 from a single disrupted asteroid (Greenwood et al., 2006). The homogeneous O isotopic composition
 286 of these pallasites provides a strong evidence for an early extensive melting event on their parent
 287 body, possibly resulting in the formation of a global magma ocean (Greenwood et al., 2006). Fe
 288 isotopes in MGPs cannot record the early stages of core segregation, because the temperatures reached
 289 during the global melting event were too high ($\gg 1300^\circ\text{C}$) to fractionate these isotopes between
 290 silicates and metallic melts. Indeed, the average Fe isotopic composition of olivines ($\delta^{56}\text{Fe} = 0.009 \pm$
 291 0.014 ‰ , $n=11$, Weyer et al., 2005) is indistinguishable from the chondritic average.

292

293 5. Conclusions

294 The marked S depletions, the siderophile element abundances and the high $\delta^{56}\text{Fe}$ values
295 displayed by ureilites point to an efficient segregation of S-rich metallic melts in the body, more likely
296 before the onset of silicate melting. Although the UPB has been disrupted, samples of its core have not
297 yet been recovered. The lack of such samples is not surprising because the number of meteorites from
298 the UPB is still limited. The fact that only one large clast of ureilitic lava is actually known
299 demonstrates that our ureilite meteorites provide only a partial sampling of the whole parent body.

300 The percolation of S-rich metallic melts has been extensively investigated in a solid matrix
301 rich in olivine, analogous to an asteroidal mantle. Metallic melts over a percolation threshold ranging
302 from about 6 vol% to 18 vol% (Yoshino et al., 2003; Bagdassarov et al., 2009; Watson and Roberts,
303 2011) can potentially create a stable interconnected network in an olivine-rich solid matrix that could
304 have allowed the early segregation of some core material. Yet the permeability of the core forming Fe-
305 FeS melts within a silicate matrix is very low, and it was inferred that percolation of these melts
306 cannot be a major process of core formation in planetesimals (Watson and Roberts, 2011). The
307 geochemical data on ureilites seem at odds with previous experiments. Certainly other factors can
308 contribute to the segregation of Fe-FeS melts and to fast core formation in planetesimals, such as shear
309 deformation of the body (Bruhn et al., 2000; Rushmer and Petford, 2011).

310

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References

- 321
322
323 Bagdassarov, N., Golabek, G.J., Solferino, G., Schmidt, M.W. (2009) Constraints on the Fe–S melt connectivity in mantle
324 silicates from electrical impedance measurements. *Phys. Earth Planet. Inter.* **177**, 139–146.
325
326 Bischoff, A. et al. (2014) Trachyandesitic magmatism in the early Solar System. *Proc. Natl. Acad. Sci. USA* **111**, 35, 12689-
327 12692.
328
329 Blichert-Toft J., Moynier F., Lee C.T.A., Telouk P., Albarede F. (2010) The early formation of the IVA iron meteorite parent
330 body. *Earth Planet. Sci. Lett.* **296**, 469-480.
331
332 Bruhn, D., Groebner, N., Kohlstedt, D.L. (2000) An interconnected network of core-forming melts produced by shear
333 deformation. *Nature* **403**, 883-886.
334
335 Cohen, B.A., Goodrich, C.A., Keil, K. (2004) Feldspathic clast populations in polymict ureilites: Stalking the missing basalts
336 from the ureilite parent body. *Geochim. Cosmochim. Acta* **68**, 4249–4266.
337
338 Clayton R.N., Mayeda T.K. (1996) Oxygen isotope studies of achondrites. *Geochim. Cosmochim. Acta* **60**, 1999-2017.
339
340 Craddock, P.R., Dauphas, N. (2011) Iron isotopic compositions of geological reference materials and chondrites. *Geostand.*
341 *Geoanal. Res.* **35**, 101–123.
- 342 Craddock, P.R., Warren, J.M., Dauphas, N. (2013) Abyssal peridotites reveal the near-chondritic Fe isotopic composition of
343 the Earth. *Earth Planet. Sci. Lett.* **365**, 63–76.
- 344 Dauphas N. et al. (2014) Magma redox and structural controls on iron isotope variations in the Earth’s mantle and crust.
345 *Earth Planet. Sci. Lett.* **398**, 127-140.
346
347 Dauphas, N., Rouxel, O. (2006) Mass spectrometry and natural variations of iron isotopes. *Mass Spectrometry Reviews* **25**,
348 515-550.
349
350 Day, J.M.D. et al. (2012) Origin of felsic achondrites Graves Nunataks 06128 and 06129, and ultramafic brachinites and
351 brachinite-like achondrites by partial melting of volatile-rich primitive parent bodies. *Geochim. Cosmochim. Acta* **81**, 94–
352 128.
353
354 Downes, H., Mittlefehldt, D.W., Kita, N.T., Valley, J.W. (2008) Evidence from polymict ureilite meteorites for a disrupted
355 and re-accreted single ureilite parent asteroid gardened by several distinct impactors. *Geochim Cosmochim Acta* **72**, 4825–
356 4844.
357
358 Goodrich, C. A., Wlotzka, F., Ross, D. K. and Bartoschewitz, R. (2006) NWA 1500: plagioclase-bearing monomict ureilite
359 or ungrouped achondrite? *Meteorit. Planet. Sci.* **41**, 925–952.
360
361 Goodrich C.A., Wilson L., van Orman J.A., Michel P. (2013a) Comment on “Parent body depth-pressure-temperature
362 relationships and the style of the ureilite anatexis” by P. H. Warren (MAPS 47:209–227) *Meteoritics & Planetary Science* **48**
363 1096-1106.
364
365 Goodrich, C. A., Ash, R. D., Van Orman, J. A., Domanik, K., McDonough, W. F. (2013b) Metallic phases and siderophile
366 elements in main group ureilites: implications for ureilite petrogenesis. *Geochim. Cosmochim. Acta* **112**, 340–373.
367
368 Goodrich C.A., Sutton S.R., Wirick S., Jercinovic M.J. (2013c) Chromium valences in ureilite olivine and implications for
369 ureilite petrogenesis. *Geochim. Cosmochim. Acta* **122**, 280-305.
370
371 Goodrich C.A., Harlow G.E., Van Orman J.A., Sutton S.R., Jercinovic M.J., Mikouchi T. (2014) Petrology of chromite in
372 ureilites: Deconvolution of primary oxidation states and secondary reduction processes. *Geochim. Cosmochim. Acta* **135**,
373 126-169.
374
375 Greenwood, R.C., Franchi, I.A., Jambon, A., Buchanan, P. (2005) Widespread magma oceans on asteroidal bodies in the
376 early solar system. *Nature* **435**, 916-918.
377
378 Greenwood, R.C., Franchi, I.A., Jambon, A., Barrat, J.A., Burbine, T.H. (2006) Oxygen isotope variation in stony-iron
379 meteorites. *Science* **313**, 1763-1765.
380
381 Halliday, A.N. (2013) Small differences in sameness. *Nature* **497**, 43-45.
382

- 383 Hin, R.C., Schmidt, M.W., Bourdon, B. (2012) Experimental evidence for the absence of iron isotope fractionation between
 384 metal and silicate liquids at 1 GPa and 1250-1300°C and its cosmochemical consequences. *Geochim. Cosmochim. Acta* **93**,
 385 164-181.
- 386
 387 Hintenberger H., Jochum K.P., Braun O., Christ P., Martin W. (1978) The antarctic meteorite Yamato 74123 – a new ureilite.
 388 *Earth Planet. Sci. Lett.* **40**, 187-193.
- 389
 390 Keil, K. (2000) Thermal alteration of asteroids: evidence from meteorites. *Planet. Space Sci.* **48**, 887-903.
- 391
 392 Keil, K. (2014) Brachinite meteorites: Partial melt residues from an FeO-rich asteroid. *Chemie der Erde* **74**, 311-329.
- 393
 394 Kruijjer, T.S. et al. (2014) Protracted core formation and rapid accretion of protoplanets. *Science* **344**, 1150-1154.
- 395
 396 Kullerud G. (1963) The Fe-Ni-S system. *Ann. Rep. Geophys. Lab.* **67**, 4055-4061.
- 397
 398 Mittlefehldt, D.W., McCoy, T.J., Goodrich, C.A., Kracher, A., (1998) Non-chondritic meteorites from asteroidal bodies. In:
 399 Papike, J.J.(Ed.), Planetary Materials. Mineralogical Society of America, Washington, DC, p.195.
- 400
 401 Moynier F, Beck P., Yin Q.Z., Ferroir T., Barrat J.A., Paniello R., Telouk P., Gillet P. (2010) Volatilization induced by
 402 impacts recorded in Zn isotope composition of ureilites. *Chem. Geol.* **276**, 374-379.
- 403
 404 Myiamoto M., Takeda H., Toyoda H. (1985) Cooling history of some Antarctic ureilites. *J Geophys. Res.* **90**, supplement,
 D116-D122.
- 405
 406 Poitrasson, F., Halliday, A.N., Lee, D.C., Levasseur, S., Teutsch, N. (2004) Iron isotope differences between Earth, Moon,
 Mars and Vesta as possible records of contrasted accretion mechanisms. *Earth Planet. Sci. Lett.* **223**, 253–266.
- 407
 408 Poitrasson, F., Roskosz, M. and Corgne, A. (2009) No iron isotope fractionation between molten alloys and silicate melt to
 409 2000 degrees C and 7.7 GPa: experimental evidence and implications for planetary differentiation and accretion. *Earth*
 410 *Planet. Sci. Lett.* **278**, 376–385.
- 411
 412 Polyakov, V. B., Clayton, R. N., Horita, J. and Mineev, S. D. Equilibrium iron isotope fractionation factors of minerals:
 413 reevaluation from the data of nuclear inelastic resonant X-ray scattering and Mössbauer spectroscopy. *Geochim. Cosmochim.*
 414 *Acta* **71**, 3833–3846 (2007).
- 415
 416 Polyakov, V. B. and Soultanov, D. M (2011) New data on equilibrium iron isotope fractionation among sulfides: constraints
 417 on mechanisms of sulfide formation in hydrothermal and igneous systems. *Geochim. Cosmochim. Acta* **75**, 1957–1974.
- 418
 419 Pringle E.A., Moynier F., Savage P.S., Badro J., Barrat J.A. (2014) Silicon isotopes in angrites and volatile loss in
 420 planetesimals. *Proc. Natl. Acad. Sci. USA*, doi: 10.1073/pnas.1418889111.
- 421
 422 Rankenburg, K., Humayun, M., Brandon, A. D., Herrin, J. S. (2008) Highly siderophile elements in ureilites. *Geochim.*
 423 *Cosmochim. Acta* **72**, 4642–4659.
- 424
 425 Rushmer T., Petford N. (2011) Microsegregation rates of liquid Fe-Ni-S metal in natural silicate-metal systems: A combined
 426 experimental and numerical study. *Geochim Geophys. Geosyst.* **12**, Q03014, doi: 10.1029/2010GC003413.
- 427
 428 Saunier G., Poitrasson F., Moine B., Gregoire M., Seddiki A. (2010) Effect of hot desert weathering on the bulk-rock iron
 429 isotope composition of L6 and H5 ordinary chondrites. *Meteorit. Planet. Sci.* **45**, 195-209.
- 430
 431 Schoenberg, R., von Blanckenburg, F. (2006) Modes of planetary-scale Fe isotope fractionation. *Earth Planet. Sci. Lett.* **252**,
 432 342–359.
- 433
 434 Schuessler, J.A., Schoenberg, R., Behrens, H., von Blanckenburg, F. (2007) The experimental calibration of the iron isotope
 435 fractionation factor between pyrrhotite and peralkaline rhyolitic melt. *Geochim. Cosmochim. Acta* **71**, 417-433.
- 436
 437 Scott, E.R.D., Taylor, G.J., Keil, K. (1993) Origin of ureilite meteorites and implications for planetary accretion. *Geophys.*
 438 *Res. Lett.* **20**, 415–418.
- 439
 440 Shahar, A. et al. (2015) Sulfur-controlled iron isotope fractionation experiments of core formation in planetary bodies.
 441 *Geochim. Cosmochim. Acta* **150**, 253-264.
- 442
 443 Singletary, S. J. and Grove, T. L. (2003) Early petrologic processes on the ureilite parent body. *Meteorit. Planet. Sci.* **38**, 95–
 444 108.
- 445
 446 Teng F.Z., Dauphas N., Huang S., Marty B. (2013) Iron isotopic systematics of oceanic basalts. *Geochim. Cosmochim. Acta*
107, 12-26.

- 447
448 Vdovykin G. P. (1970) Ureilites. *Space Sci. Rev.* **10**, 483–510.
449
- 450 Wang, K. *et al.* (2012) Iron isotope fractionation in planetary crusts. *Geochim. Cosmochim. Acta* **89**, 31–45.
451
- 452 Wang, K. *et al.* (2013) Homogeneous distribution of Fe isotopes in the early solar nebula. *Meteorit. Planet. Sci.* **48**, 354–364.
453
- 454 Wang, K. *et al.* (2014) Iron isotope fractionation during sulfide-rich felsic partial melting in early planetesimals. *Earth Planet. Sci. Lett.* **392**, 124–132.
455
- 456 Wänke H., Baddenhausen H., Spettel B., Teschke F., Quijano-Rico M., Dreibus G., Palme H. (1972) The chemistry of , 572-
457 590Haverö ureilite. *Meteoritics* **7**, 579-590.
458
- 459 Warren P.H. (2011) Stable isotopes and the noncarbonaceous derivation of ureilites, in common with nearly all differentiated
460 planetary materials. *Geochim. Cosmochim. Acta* **75**, 6912-6926.
461
- 462 Warren P. H. (2012) Parent body depth-pressure-temperature relationships and the style of the ureilite anatexis. *Meteoritics
& Planetary Science* **47**, 209–227.
463
- 464 Warren, P. H. and Rubin, A. E. (2010) Pyroxene-selective impact smelting in ureilites. *Geochim. Cosmochim. Acta* **74**, 5109–
465 5133.
466
- 467 Warren, P.H., Ulff-Moller, F., Huber, H., Kallemeyn, G.W. (2006) Siderophile geochemistry of ureilite: a record of early
468 stages of planetesimal core formation. *Geochim. Cosmochim. Acta* **70**, 2104-2126.
469
- 470 Wasson, J.T., Kallemeyn, G.W. (1988) Compositions of chondrites. *Phil. Trans. R. Soc. London A* **325**, 535-544.
471
- 472 Watson, H.C., Roberts, J.J. (2011) Connectivity of core forming melts: Experimental constraints from electrical conductivity
473 and X-ray tomography. *Phys. Earth Planet. Inter.* **186**, 172-182.
474
- 475 Weyer, S. *et al.* (2005) Iron isotope fractionation during planetary differentiation. *Earth Planet. Sci. Lett.* **240**, 251–264.
476
- 477 Yanai K., Kojma H., Haramura H. (1995) Catalog of the Antarctic meteorites. National Institute of Polar Research, Tokyo,
478 230 p.
479
- 480 Yoshino, T., Walter, M.J., Katsura, T. (2003) Core formation in planetesimals triggered by permeable flow. *Nature* **422**, 154–
481 157.
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484 Table 1. Details of meteorite samples studied. Olivine compositions are from this study except D
 485 (Downes et al., 2008), SG (Singletary and Grove, 2004), G (Goodrich et al., 2006), and WR (Warren
 486 and Rubin, 2010).

	Source	# or split	Olivine core Fo %	bulk rock Mg#
<i>Fall</i>				
Novo Urei	Saint Louis		78.5 ^{SG}	
<i>Hot desert finds</i>				
El Gouanem	ENS Lyon		80.7 ^{SG}	
NWA 2236	NIPR	,41	96.8	91.8
NWA 4471	JAB		78.1	70.2
NWA 4509	ENS Lyon		78.5	
NWA 4511	ENS Lyon		77.8	77.2
NWA 4512	ENS Lyon		78.3	
NWA 4513	ENS Lyon		90.5	
NWA 4516	ENS Lyon		81.3	76.9
NWA 5555	JAB		90.8	87.3
NWA 5602	JAB		79.0	76.5
NWA 5884	JAB		78.6	75.0
NWA 6056	JAB		84.8	81.4
NWA 7349	JAB		76.5	75.4
NWA 7630	JAB		79.1	73.4
NWA 7686	JAB		91.0	84.3
NWA 7880	JAB		78.6	77.7
NWA 8049	JAB		84.3	81.3
<i>Antarctic finds</i>				
A-881931	NIPR	,65	78.7	76.5
ALHA 77257	NIPR	,104	86.1 ^D	83.7
ALH 82130	MWG	,43	95.2 ^D	89.9
EET 83225	MWG	,37	88.3 ^D	85.5
LAP 03587	MWG	,10	74.7 ^{WR}	74.1
LAR 04315	MWG	,46	81.9 ^{WR}	79.1
MET 01085	MWG	,23	no olivine	84.9
Y-791538	NIPR	,126	91.3 ^D	87.1
Y-981810	NIPR	,88	78.3	74.4
<i>Other finds</i>				
Kenna	Saint Louis		78.0 ^G	
Goalpara	Saint Louis		76.6 ^D	
<i>Ureilitic trachyandesite (Almahata Sitta fall)</i>				
ALM-A	Münster		no olivine	

488 Table 2. Olivine core compositions of the samples analyzed during the course of this study (oxides in
 489 wt%, Fe/Mn atomic).

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	lab.	n	SiO ₂	Cr ₂ O ₃	FeO	MnO	MgO	CaO	total	Fo%	Fe/Mn
A-881931	NIPR	13	38.07	0.49	19.72	0.40	40.78	0.29	99.75	78.7	48.1
NWA2236	NIPR	28	42.38	0.38	3.21	0.45	53.91	0.28	100.64	96.8	7.0
NWA 4471	SCMO	10	38.75	0.49	19.92	0.39	39.92	0.29	99.76	78.1	50.1
NWA 4509	SCMO	10	39.06	0.75	19.73	0.41	40.49	0.37	100.80	78.5	48.0
NWA 4511	SCMO	10	38.80	0.54	20.32	0.40	40.03	0.31	100.40	77.8	50.7
NWA 4512	SCMO	10	39.47	0.71	19.75	0.40	39.95	0.37	100.67	78.3	48.2
NWA 4513	SCMO	10	40.82	0.59	9.08	0.43	48.52	0.30	99.74	90.5	20.8
NWA 4516	SCMO	10	39.19	0.67	17.38	0.41	42.28	0.32	100.26	81.3	41.5
NWA5555	SCMO	10	41.07	0.62	8.89	0.45	49.36	0.31	100.72	90.8	19.5
NWA 5602	SCMO	10	38.60	0.72	19.32	0.41	40.88	0.35	100.28	79.0	46.3
NWA5884	SCMO	10	39.32	0.70	19.36	0.40	39.92	0.32	100.06	78.6	47.4
NWA6056	SCMO	10	40.26	0.75	14.27	0.44	44.73	0.39	100.87	84.8	31.8
NWA 7349	SCMO	10	38.54	0.39	21.55	0.41	39.29	0.23	100.41	76.5	51.3
NWA 7630	SCMO	10	38.89	0.74	19.26	0.41	40.78	0.33	100.40	79.1	46.6
NWA 7686	SCMO	10	40.88	0.64	8.65	0.45	49.16	0.34	100.15	91.0	19.1
NWA 7880	SCMO	10	38.82	0.74	19.69	0.40	40.58	0.28	100.54	78.6	49.2
NWA 8049	SCMO	10	39.85	0.71	14.66	0.42	44.11	0.37	100.11	84.3	34.5
Y-981810	NIPR	18	38.64	0.55	20.05	0.39	40.54	0.35	100.51	78.3	50.2

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494 Table 3. Iron isotopic compositions of international standards obtained during the course of this study
 495 in Saint Louis and in Plouzané, and compared with literature values. (N# = number of runs of the same
 496 solution).

	N#	$\delta^{56}\text{Fe}$	95% C.I.	$\delta^{57}\text{Fe}$	95% C.I.
BIR1					
Plouzané, average (n=6)		0.073	±0.017	0.116	±0.013
Wang et al. (2012)		0.043	±0.016	0.058	±0.026
Craddock & Dauphas (2011)		0.053	±0.015	0.087	±0.023
BHVO2					
Plouzané, average (n=14)		0.117	±0.011	0.181	±0.023
Wang et al. (2012)		0.102	±0.012	0.159	±0.018
Craddock & Dauphas (2011)		0.114	±0.011	0.174	±0.016
Allende USNM3529					
Plouzané #1	5	0.015	±0.015	0.014	±0.069
Plouzané #2	5	0.021	±0.035	-0.019	±0.039
Plouzané, average (n=2)		0.018		-0.003	
Craddock & Dauphas (2011)		-0.007	±0.012	0.003	±0.019
AC-E					
Plouzané #1	5	0.316	±0.025	0.501	±0.065
Wang et al. (2012)		0.313	±0.018	0.457	±0.028
Craddock & Dauphas (2011)		0.320	±0.010	0.478	±0.015

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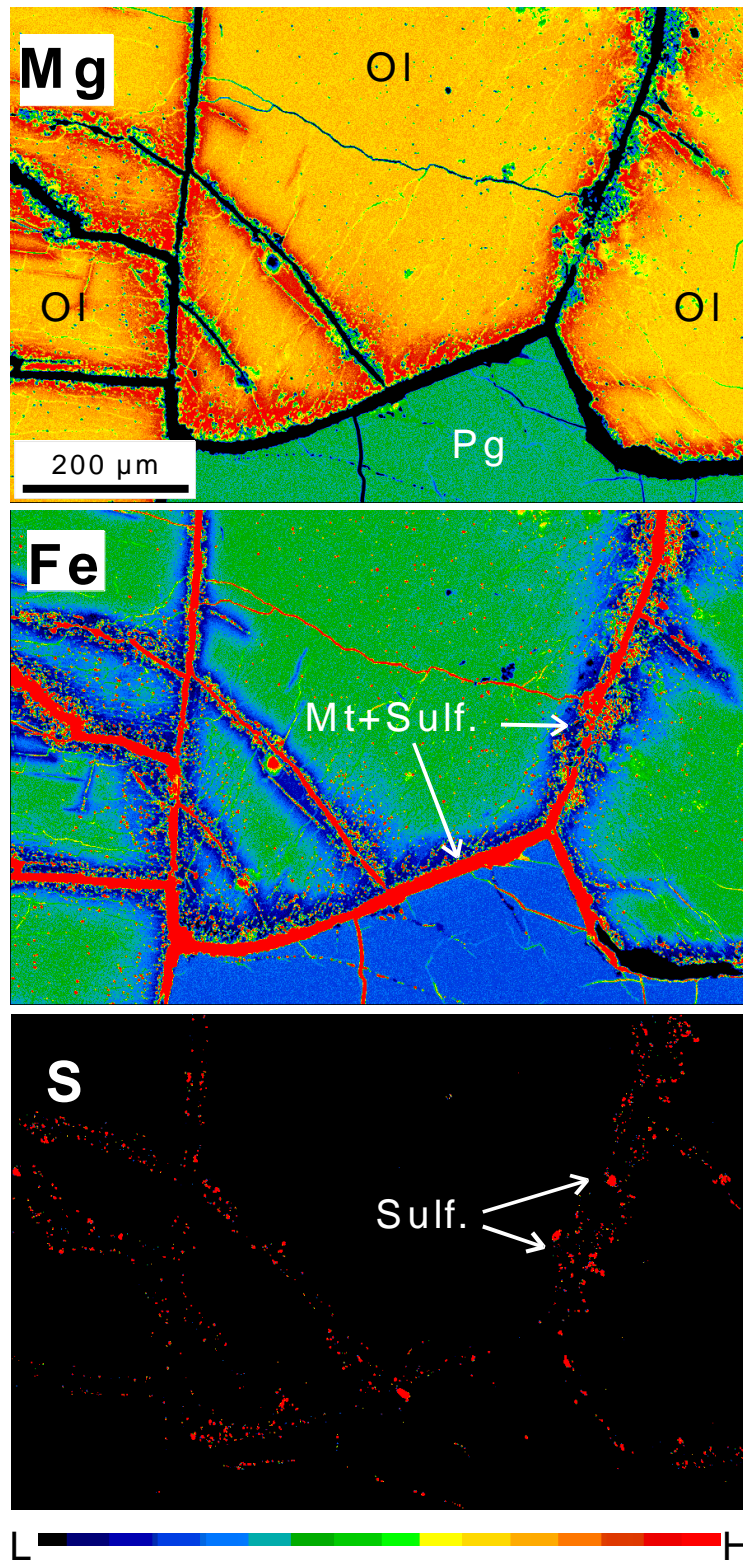
500 Table 4. Iron isotopic compositions of unbrecciated and trachyandesitic ureilites.

	lab.	N#	$\delta^{56}\text{Fe}$	95% C.I.	$\delta^{57}\text{Fe}$	95% C.I.	
<i>Fall</i>							
	Novo Urei #1	St Louis	9	0.024	± 0.030	0.027	± 0.050
	Novo Urei #2	St Louis	9	0.039	± 0.038	0.084	± 0.057
<i>Northwest Africa finds</i>							
	El Gouanem	St Louis	9	0.068	± 0.028	0.085	± 0.052
	NWA 2236	Plouzané	6	0.076	± 0.017	0.129	± 0.047
	NWA 4471	Plouzané	8	0.086	± 0.021	0.133	± 0.033
	NWA 4509	St Louis	9	0.024	± 0.033	0.060	± 0.058
	NWA 4511 #1	Plouzané	8	0.051	± 0.032	0.073	± 0.052
	NWA 4511 #2	St Louis	8	0.023	± 0.032	0.062	± 0.059
	NWA 4512	St Louis	8	0.046	± 0.032	0.082	± 0.059
	NWA 4513	St Louis	9	0.054	± 0.038	0.098	± 0.057
	NWA 4516	Plouzané	9	0.047	± 0.030	0.073	± 0.044
	NWA 5555	Plouzané	7	0.053	± 0.031	0.081	± 0.047
	NWA 5602	Plouzané	10	0.043	± 0.029	0.065	± 0.049
	NWA 5884	Plouzané	7	0.057	± 0.017	0.087	± 0.028
	NWA 6056 #1	Plouzané	5	0.040	± 0.025	0.027	± 0.027
	NWA 6056 #2	Plouzané	7	0.054	± 0.021	0.080	± 0.035
	NWA 7349	Plouzané	8	0.059	± 0.018	0.094	± 0.027
	NWA 7630	Plouzané	7	0.081	± 0.026	0.124	± 0.037
	NWA 7686 #1	Plouzané	5	0.090	± 0.017	0.117	± 0.066
	NWA 7686 #2	Plouzané	7	0.067	± 0.029	0.103	± 0.047
	NWA 7880	Plouzané	6	0.040	± 0.022	0.063	± 0.030
	NWA 8049	Plouzané	7	0.067	± 0.031	0.109	± 0.055
<i>Antarctic finds</i>							
	A-881931	Plouzané	5	0.109	± 0.039	0.209	± 0.062
	ALHA 77257	Plouzané	6	0.058	± 0.030	0.079	± 0.057
	ALH 82130	Plouzané	8	0.044	± 0.034	0.053	± 0.045
	EET 83225	Plouzané	6	0.035	± 0.026	0.038	± 0.067
	LAP 03587 #1	Plouzané	7	0.035	± 0.030	0.075	± 0.016
	LAP 03587 #2	St Louis	9	0.055	± 0.033	0.082	± 0.058
	LAR 04315	Plouzané	6	0.059	± 0.029	0.088	± 0.057
	MET 01085	Plouzané	6	0.023	± 0.023	0.036	± 0.055
	Y-791538	Plouzané	6	0.071	± 0.018	0.121	± 0.045
	Y-981810	Plouzané	5	0.077	± 0.015	0.125	± 0.073
<i>Other finds</i>							
	Kenna	St Louis	9	0.009	± 0.030	0.030	± 0.049
	Goalpara #1	St Louis	9	0.048	± 0.030	0.095	± 0.049
	Goalpara #2	St Louis	9	0.056	± 0.038	0.092	± 0.057
<i>Ureilitic trachyandesite</i>							
	ALM-A #1	Plouzané	5	0.095	± 0.032	0.155	± 0.066
	ALM-A #2	Plouzané	4	0.096	± 0.019	0.107	± 0.045
	ALM-A #3	Plouzané	8	0.105	± 0.023	0.159	± 0.038

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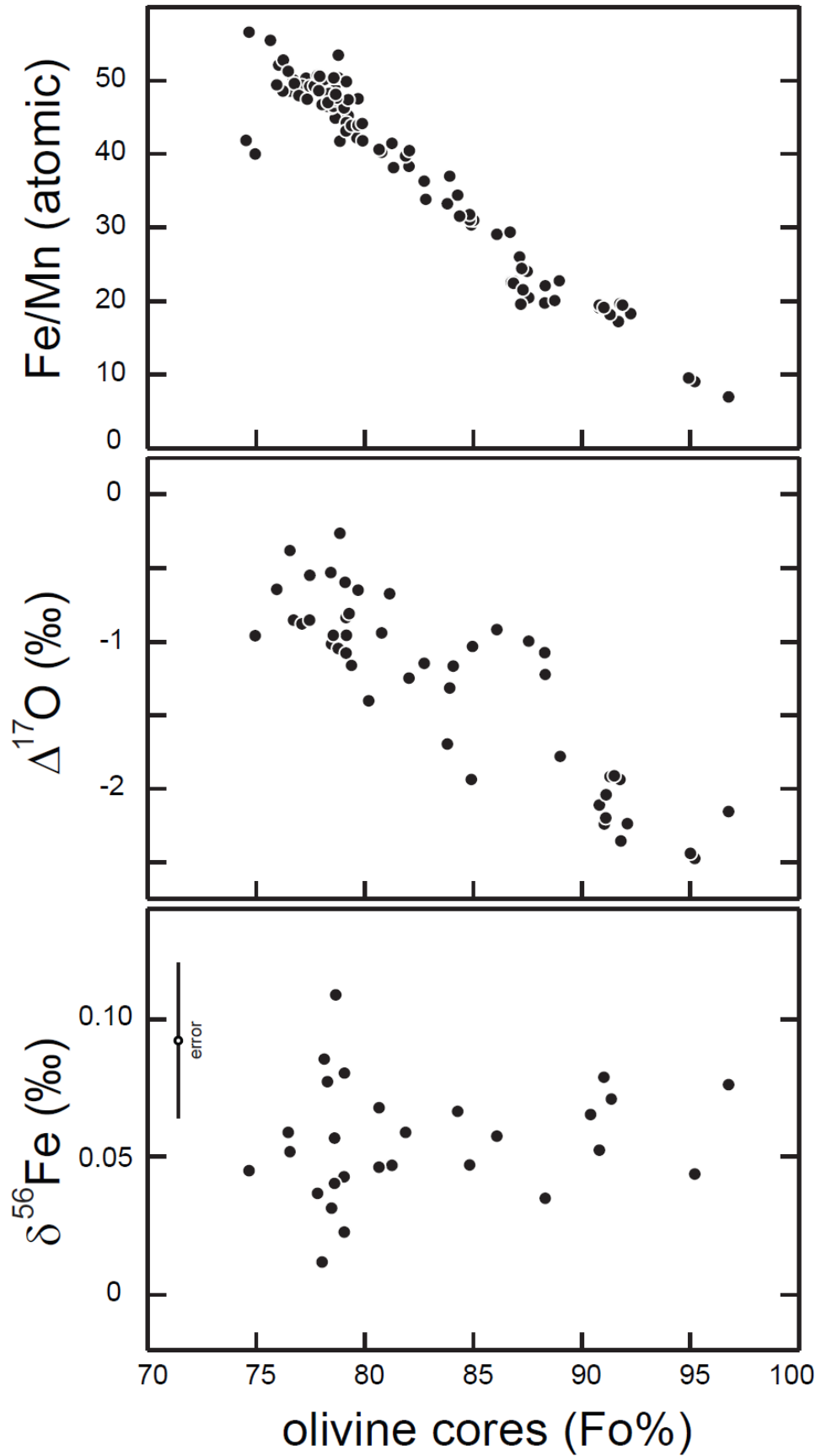
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Figure 1. Maps of Mg, Fe and S of Northwest Africa 8049, a typical ureilite, showing the zoning of the olivine grains (Ol), the homogeneity of the pigeonite (Pg), and the repartition of metal (Mt - mainly interstitial and in tiny inclusions into olivine) and sulfides (Sulf.).



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Figure 2. Plots of molar Fe/Mn ratios in olivine cores, $\Delta^{17}\text{O}$ (Clayton and Mayeda, 1996) and $\delta^{56}\text{Fe}$ in the bulk rocks vs. the composition of the olivine cores (data from Table 2, and mainly from Downes et al. (2008), Singletary and Grove (2003), Goodrich et al. (2006, 2014) and references therein).

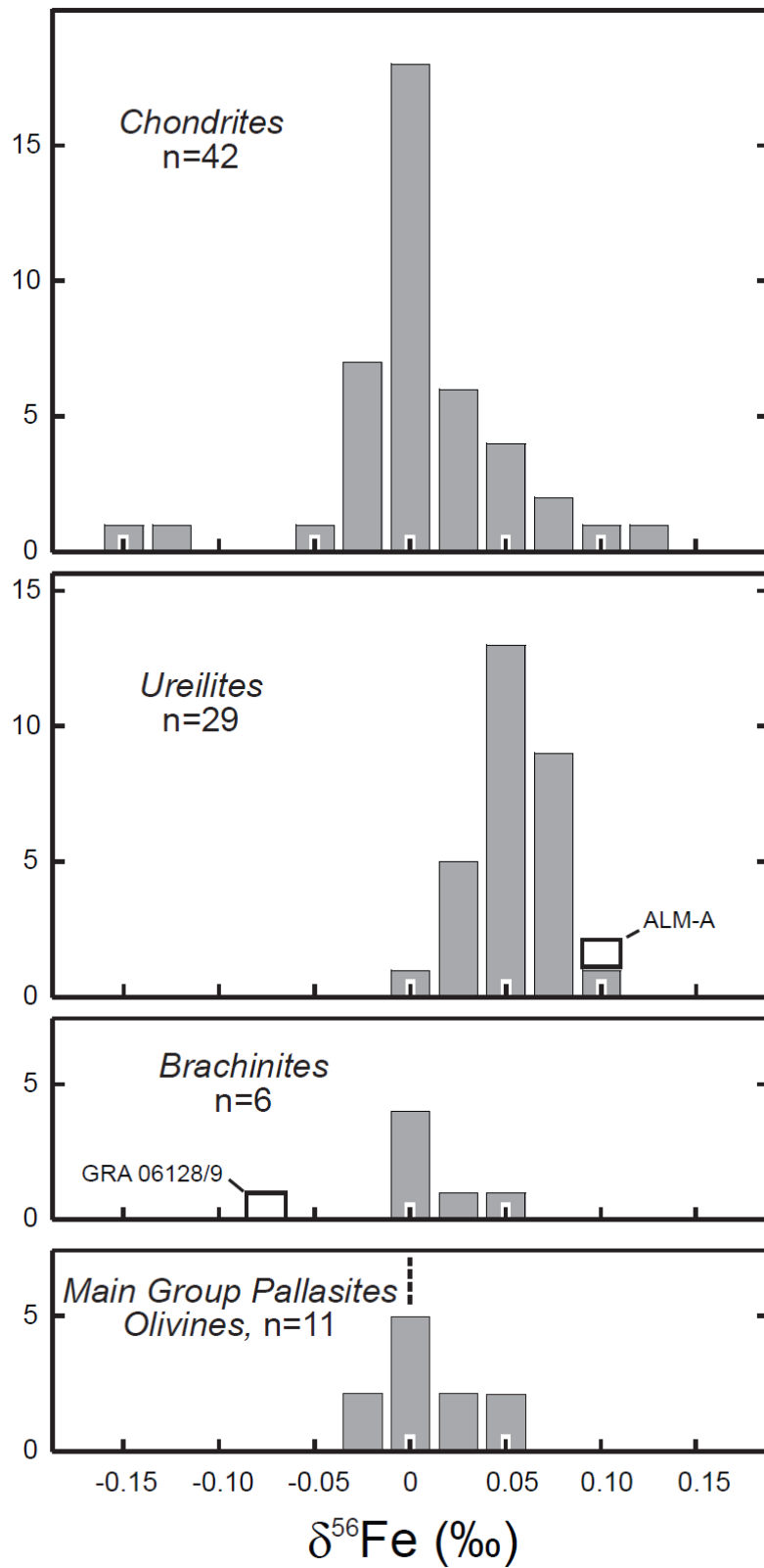
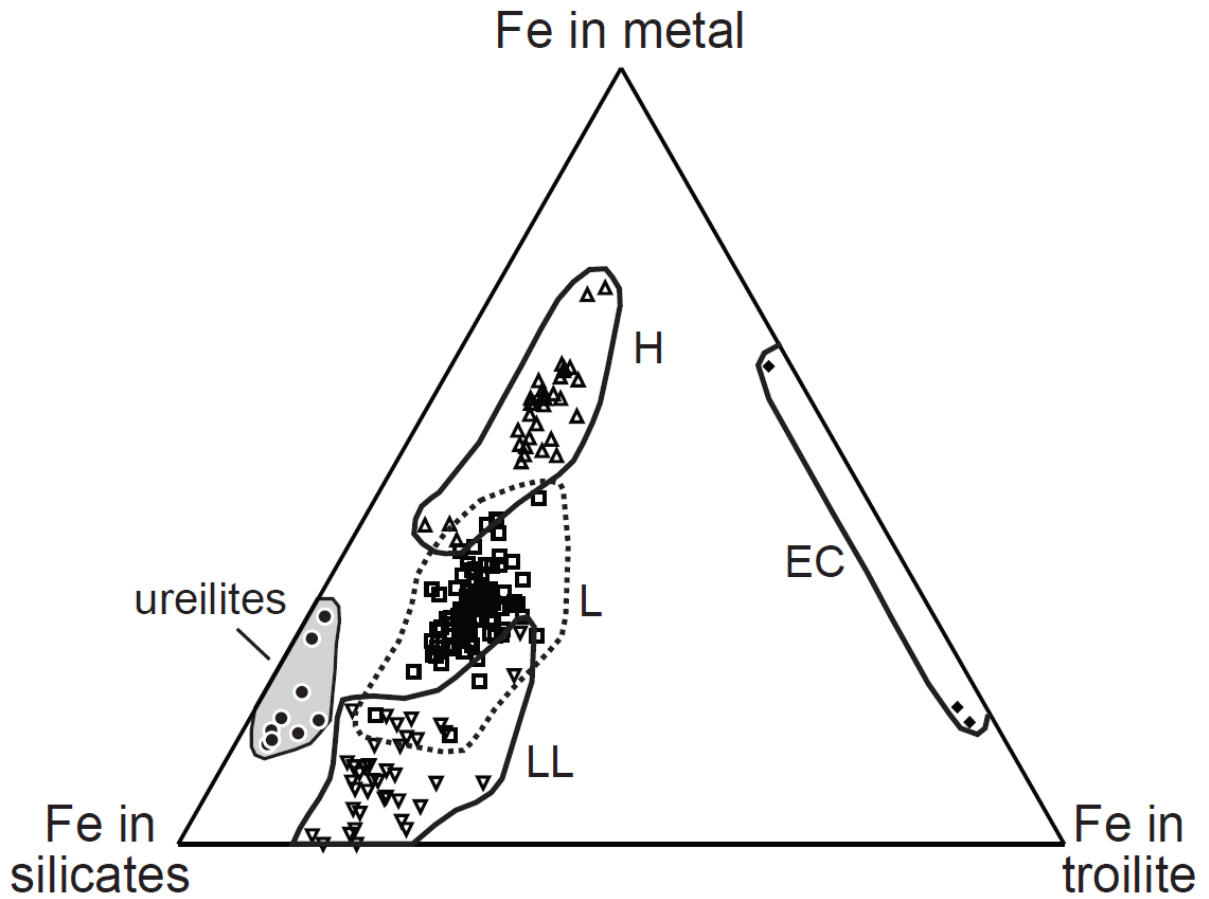
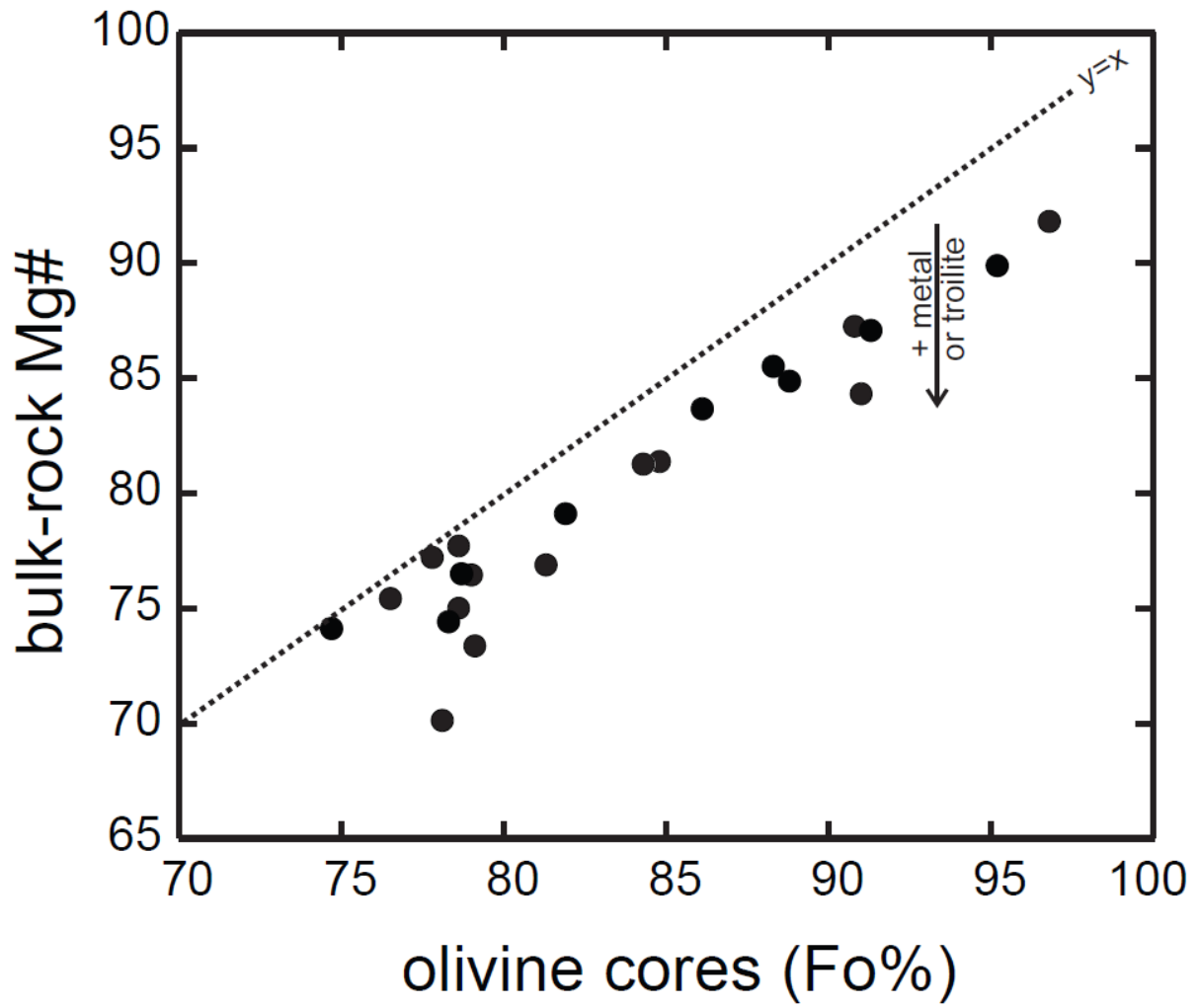
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Figure 3. Iron isotope frequency distributions of chondrites (Craddock et al., 2013), ureilites and ALM-A, a ureilitic lava (this work), brachinites and GRA 06128/9, a brachinitic melt (Wang et al., 2014), and olivines from main group pallasites (Weyer et al., 2005).



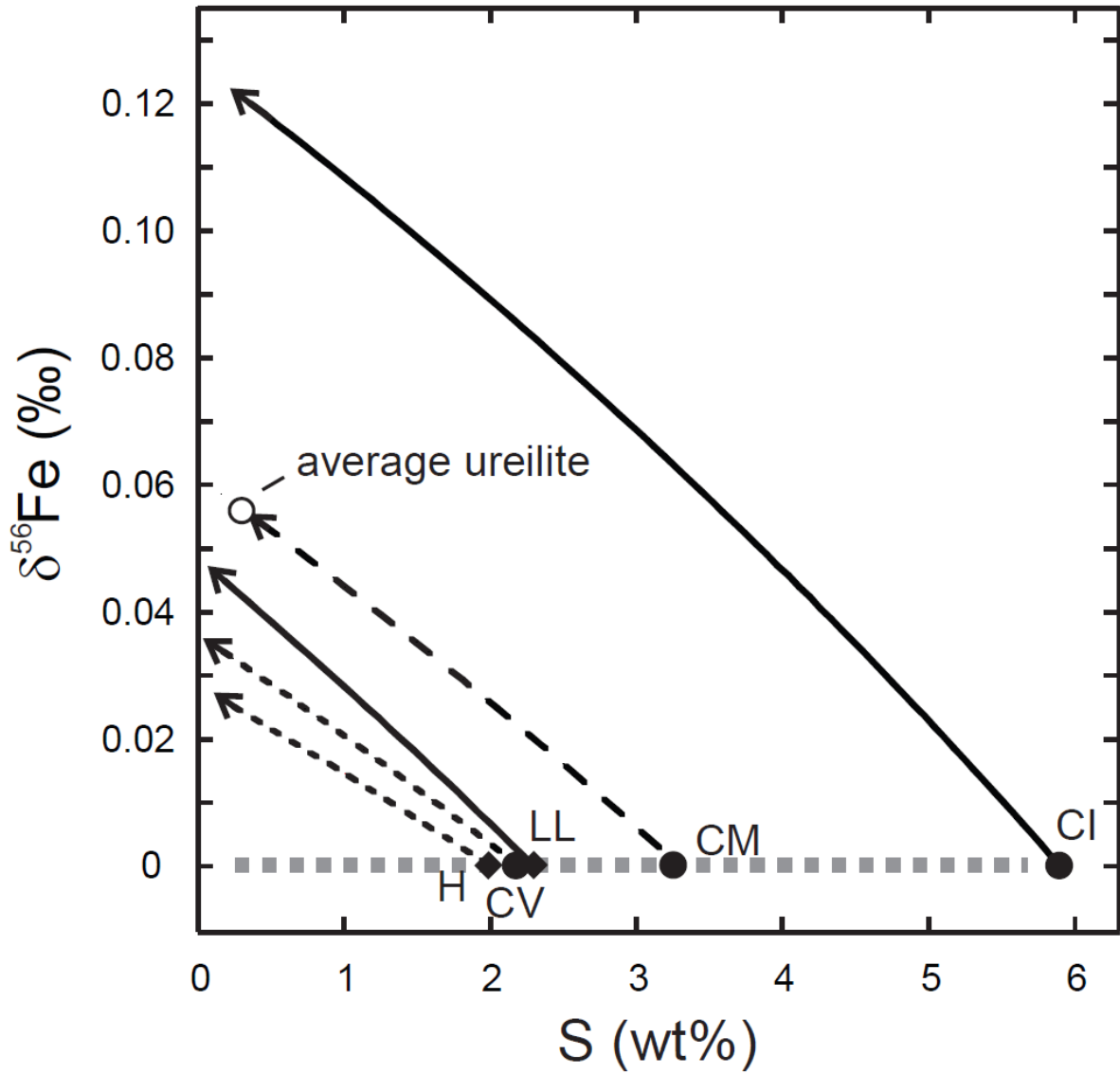
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Figure 4. Ternary diagram showing the proportions of Fe hosted in the silicates, in metal and in sulfides for ureilites and non-carbonaceous chondrites (the proportions were estimated from the analyses compiled by Yanai et al., 1995).



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Figure 5. Bulk-rock Mg# values vs. olivine-core compositions for the ureilites analyzed in Plouzané.



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Figure 6. Evolution of the isotopic compositions of residual lithologies as a function of the extraction of a Fe-FeS melt with a eutectic composition. Starting assemblages have a $\delta^{56}\text{Fe} = 0$ ‰ and the Fe and S contents of selected chondritic types (Wasson and Kallemeyn, 1988). The S abundance in the average ureilite is from Warren et al. (2006). A protolith with S and Fe abundances similar to average CM can account for the ureilite features, but this solution is not unique. Notice that the protolith of the ureilites is not a known carbonaceous chondrite (Warren, 2011).