1	High spatial resolution measurements of iron isotopes in pyrites by
2	SIMS using the new Hyperion-II Radio-Frequency Plasma source.
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27	Key words: Iron isotopes; Ion microprobe; Pyrites.
28	Abbreviations: SIMS: Secondary Ions Mass Spectrometry; MC-ICP-MS: Multi-collector
29	inductively coupled plasma mass spectrometry; IMF: Instrumental Mass Fractionation; SD:
30	Standard Deviation; SE: Standard Error.
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32 Abstract

Rationale: Iron isotopic signatures in pyrites are considered as a good proxy to reconstruct paleoenvironmental and local redox conditions. However, the investigation of micro-pyrites less than 20µm size has been limited so far by analytical techniques. The development of the new brighter radio-frequency plasma ion source (Hyperion-II source) enhances the spatial resolution by increasing the beam density 10 times compared to the Duoplasmatron source.

38 Methods: Here we present high spatial resolution measurements of iron isotopes in pyrites 39 using a 3nA-3µm primary ¹⁶O⁻ beam on two ion microprobes Cameca IMS 1280-HR2 at 40 CRPG-IPNT (France) and at SwissSIMS (Switzerland) equipped with Hyperion sources. We 41 tested analytical effects, such as topography and crystal orientation that could induce 42 analytical biases perceptible through variations of the Instrumental Mass Fractionation (IMF). **Results**: The δ^{56} Fe reproducibility for the Balmat pyrite standard is $\pm 0.25\%$ (2SD, standard 43 44 deviation) and the typical individual internal error is $\pm 0.10\%$ (2SE, standard error). The 45 sensitivity on ⁵⁶Fe⁺ was 1.2x10⁷cps/nA/ppm or better. Tests on Balmat pyrites revealed that 46 neither the crystal orientation nor channeling effects seem to significantly influence the IMF. 47 Different pyrite standards (Balmat and SpainCR) were used to test the accuracy of the 48 measurements. Indium mounts must be carefully prepared with sample topography $< 2\mu m$, 49 which was checked using an interferometric microscope. Such a topography is negligible for 50 introducing change in the IMF. This new source increases the spatial resolution while 51 maintaining the high precision of analyses and the overall stability of the measurements 52 compared to the Duoplasmatron source.

53 **Conclusions**: We developed a reliable method to perform accurate and high-resolution 54 measurements of micrometric pyrites. The investigation of sedimentary micro-pyrites will 55 improve our understanding of the processes and environmental conditions during pyrite 56 precipitation, including contribution of primary (microbial activities or abiotic reactions) and 57 secondary (diagenesis and/or hydrothermal fluid circulation) signatures.

59 **1. Introduction**

60 Iron stable isotope geochemistry has been developed rapidly over the last 15 years, particularly because iron is a ubiquitous element that occurs in three oxidation states: Fe⁰, 61 Fe^{2+} and Fe^{3+} . The redox state affects iron isotope fractionation of the four stable isotopes, 62 ⁵⁴Fe (5.80%), ⁵⁶Fe (91.72%), ⁵⁷Fe (2.20%) and ⁵⁸Fe (0.28%), following the mass- and 63 temperature-dependent fractionation laws¹⁻⁴. The iron isotopic composition is reported using 64 the delta (δ) notation (in ∞) defined as deviations of the measured ⁵⁶Fe/⁵⁴Fe or ⁵⁷Fe/⁵⁴Fe 65 ratios of the sample relative to the international standard IRMM-014 (56Fe/54Fe value of 66 15.6986 and 57 Fe/ 54 Fe=0.3626) 5,6 : 67

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$$\delta^{5x} Fe = \left[\frac{\left(\frac{5^{x} Fe}{5^{4} Fe}_{sample}\right)}{\left(\frac{5^{x} Fe}{5^{4} Fe}_{IRMM014}\right)} - 1 \right]$$

where x is either 6 or 7. Accuracy and precision were determined by the analysis of in-house
pyrite standards, Balmat and SpainCR (detailed in section 2.1).

72 The difference of Fe isotopic compositions between 2 species defines the isotopic 73 fractionation. Iron isotope systematics is used in numerous fundamental fields, such as in 74 cosmochemistry and igneous petrology that focus on the accretion of planetary bodies, 75 magmatic differentiation, and diffusion during crystal growth^{5,7-11}, as well as in 76 environmental geochemistry for understanding the past and modern redox marine cycle^{5,12-17}. 77 Iron is also a major element in numerous biological reactions leading to diagnostic isotopic 78 fractionations and thus, can be a good proxy for biosignature recognition. The microbial iron 79 cycle is controlled by Dissimilatory Iron Reduction (DIR) and bacterial iron oxidation leading to precipitation of diverse Fe(II)-bearing biominerals, including iron sulfides^{18,19}. 80 81 However, pyrites can also be formed through abiotic reactions. Both biological and abiotic pathways are associated with large mass-dependent Fe isotopic fractionations²⁰⁻³³. 82 Consequently, biologically-precipitated pyrites record δ^{56} Fe variations up to $6\%^{20,22-24}$, 83 encompassing the entire range of terrestrial Fe isotopic fractionation (see e.g., Johnson et al,¹³ 84 85 and Beard and Johnson,³⁴ for reviews). The formation of sedimentary pyrites is a complex, multi-stage process that includes dissolution of FeS_m precursors such as mackinawite (FeS) 86 87 or greigite (Fe₂S₃) ^{e.g. 35,36}. Importantly, biologically-mediated sulfides are typically very small, <1µm in size, as illustrated by FeS precipitates in microbial sulfate reducing bacteria 88 89 cultures (reviewed in Rickard³⁷). The dissolution of those FeS precursors leads to the

90 precipitation of pyrites ranging from <1 to tens of micrometers in size (especially for 91 sedimentary pyrite framboids)³⁸, which induce analytical challenges for isotopic 92 measurements in a single pyrite grain.

93 Investigation of iron isotope variability started with application of Thermal Ionization 94 Mass Spectrometry (TIMS) and was subsequently followed by Multi Collection Inductively 95 Coupled Plasma Mass Spectrometry (MC-ICPMS) yielding higher sensitivity and higher 96 precision measurements. As a result, the application of MC-ICPMS method on igneous rocks 97 allowed the recognition of small iron isotope variations associated with high temperature fractionation processes^{8,32,33}. However, only few studies focused on microscale isotope 98 99 variations in biogeochemistry. For instance, depending of the technique, strong contrasts in 100 Fe isotope compositions have been documented in pyrites from the 2.7 Ga Tumbiana 101 Formation (Pilbara craton, Western Australia). Bulk rock analyses produced a narrow range of δ^{56} Fe values of -0.02 ±0.26‰ (2SD, standard deviation)¹⁵, while *in situ* analyses by LA-102 103 MC-ICPMS (Laser Ablation Multi Collector Inductively Coupled Plasma Mass Spectrometry) yielded a large range of δ^{56} Fe values, from -2.9‰ to +1.5‰^{39,40}. Similarly, in 104 105 Archean pyrite nodules (2.7 Ga shale from Bubi Greenstone Belt, Zimbabwe), bulk δ^{56} Fe values were around -1.4‰ whereas in situ measurements by Secondary Ion Mass 106 Spectrometry (SIMS) described rim to core profiles from +0.5% to $-2\%^{16}$. These two 107 108 examples strongly support the importance of combining whole rocks and micrometer scale in 109 *situ* analysis to constrain primary and/or secondary signals recorded in a pyrite.

110 Available on the market since 2015, the new radio-frequency oxygen ion source on SIMS instruments, the Hyperion-II, has the primary beam current density improved 10 times 111 compared to the usual Duoplasmatron oxygen source⁴¹. Characteristics of the primary beam 112 and secondary transmission are documented in Liu et al.⁴¹. For a given primary beam 113 114 intensity, the improvement of current density increases spatial resolution while maintaining 115 analytical precision compared to the previous ¹⁶O⁻ source performance. The improvement of 116 the secondary ion transmission, higher precision and reproducibility measurements have been reported for Mg^{41,42} and Si⁴³ isotope systems. For example, an external reproducibility of 117 $\pm 0.2\%$ (2SD) on Δ^{26} Mg* (independent mass fractionation) is achieved with a primary beam 118 size of $3-4\mu m^{41}$, much smaller compared to $15\mu m$ with the Duoplasmatron source⁴⁴. The new 119 120 source has also higher sensitivity per unit area for Pb isotopes with a 10nA primary beam, 121 allowing the precision of U-Pb dating measurements of zircons to be enhanced⁴¹. Therefore, 122 one of the main advantages of the Hyperion source is the improved spatial resolution 123 achieved without a loss in precision required for resolving biogeochemical processes.

Previously, the expected precision of $\pm 0.2\%$ (2SD) for iron isotope analysis was obtained with a 10nA primary intensity delivered by the Duoplasmatron, focused into a ~15µm spot to generate enough secondary ion intensity to be collected by Multi Collection Faraday Cups (MCFC). However, such spot sizes limited the investigation of Fe isotopes in micrometric grains, whereas the Hyperion source opens new possibilities for the studies of biochemical processes.

In order to explore micrometric scale variations of iron isotope in pyrites, we have adapted a previously described analytical procedure by using the Hyperion source, combining the reduction of the spot size for a given intensity, and maintaining a precision of 0.2‰ (2SD). This protocol has been tested on two different SIMS instruments and will open a new analytical field to study iron isotope microscale variabilities in sedimentary pyrites.

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136 **2. Experimental**

SIMS measurements were carried out using the Cameca[®] IMS 1280-HR2 ion probes
at CRPG-IPNT (Nancy, France) and SwissSIMS (University of Lausanne, Switzerland), both
equipped with the new Hyperion-II Radio frequency source. We performed four Fe isotope
sessions at CRPG-IPNT (February 2018, April 2018, July 2018 and September 2020) and
four sessions at SwissSIMS (July 2019, January 2020, March 2020 and June 2020).

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143 *2.1. Standards*

The stability of both instruments was monitored by running multiple measurements of 144 145 in-house pyrite standards, either Balmat or SpainCr pyrites, that were also used to correct a possible instrumental drift. The major element composition of Balmat pyrite is described in 146 Marin-Carbonne et al.⁴⁵ and it contains 46.5wt.% of iron and 53.6wt.% of S. This pyrite 147 standard is extensively used for SIMS Fe and S isotopes analyses^{16,45-47}. The major and trace 148 149 element compositions of SpainCR pyrite was determined using a JEOL JXA-8530F electron 150 microprobe at the University of Lausanne and data are available in supporting information 151 A. The analytical parameters are described in supporting information B. SpainCR pyrite 152 grains contain 46.6wt.% of Fe and 53.1wt.% of S. Co, Mn, Cr, Zn and Cu contents are below 153 the detection limit. Pb, Ti and Ni contents are 1112ppm, 1003ppm and 354ppm, respectively. The SpainCR standard was previously used for SIMS S isotope analysis^{47,48}. The Fe isotopic 154 155 composition of SpainCR pyrite was determined at CRPG. Chemistry for Fe purification, and Fe isotopic analysis using the NeptunePlus MC-ICP-MS are detailed elsewhere^{45,49,50}. In 156 157 order to evaluate the homogeneity of Fe isotopic composition of SpainCR pyrite standard,

158 core and rim separates have been chemically processed and analyzed. The bulk Fe isotope 159 measurement sequence follows the sample/standard bracketing method, with IRMM-014 Fe as the normalizing standard. We obtained similar δ^{56} Fe (and δ^{57} Fe) values of +0.516 ±0.05‰ 160 $(+0.804 \pm 0.03\%)$ (2SD) and $+0.521 \pm 0.02\%$ ($+0.795 \pm 0.01\%$) (2SD) for core and rim, 161 respectively, for n= 3 replicates of each. This gives a δ^{56} Femean reference value of 0.52 162 163 $\pm 0.03\%$ for this highly homogeneous SpainCR pyrite standard. Data accuracy and 2SD 164 reproducibility are evaluated by replicate analyses of geostandards, with values of δ^{56} Fe = $\pm 0.648 \pm 0.129$ ‰ and δ^{57} Fe = $\pm 0.960 \pm 0.163$ ‰ (n=3) for IF-G (BIF Greenland), and δ^{56} Fe = 165 $+0.098 \pm 0.033\%$ and δ^{57} Fe = $+0.143 \pm 0.057\%$ (n=15) for BIR-1 (USGS Iceland basalt) 166 167 during the course of this study. These values are within the range of the reported reference 168 values⁵. The homogeneity of SpainCR standard was tested by SIMS by measuring three 169 different grains prepared in the same mount (July 2018 session at CRPG-IPNT). Based on 68 170 measurements, the external reproducibility was $\pm 0.28\%$ (2SD) (Figure S1, supporting 171 information **B**).

As no chromium was detected in the pyrite reference materials, Russie magnetite standard was used to evaluate the degree of ⁵⁴Cr interference on ⁵⁴Fe, which was necessary for the natural samples due to the presence of Cr. Details are given in section 2.5.

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2.2. Sample preparation

177 Standard grains were embedded in epoxy and polished with $0.5\mu m$ diamond paste. 178 Those grains were carefully removed from epoxy and pressed into 1-inch indium mount. 179 White light interferometric microscope (Bruker Contour GTK at University of Lausanne) was 180 used to measure the sample topography. The relief across the analyzed surface was below 181 $5\mu m^{51}$. Samples were coated with a 35nm thick gold film to ensure the conductivity between 182 the sample surface and the SIMS holder.

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2.3. Electron backscatter diffraction (EBSD):

In order to check for the influence of crystallographic orientations on SIMS measurements, three Balmat pyrite grains were removed from the Indium mount, embedded in epoxy resin with the analyzed mineral surfaces facing upwards. To eliminate any residual surface damage, the mount was further polished for 40 min using the combined chemical and mechanical effect of an alkaline (pH=9.8) suspension of colloidal silica (0.05 μ m). The sample was not carbon-coated before electron backscatter diffraction (EBSD) analysis in 191 order to maximize the detected signal. The crystallographic orientation patterns were 192 acquired at the University of Lausanne, Switzerland, using a Tescan Mira II LMU field 193 emission-scanning electron microscope (FE-SEM) equipped with the Symmetry detector and 194 the Aztec 4.2 software package, both released by Oxford Instruments®. Acquisition 195 parameters included an acceleration voltage of 20 kV, a probe current of 1.1 nA, a working 196 distance of 23 mm, and a 70° tilt of the sample surface with respect to the horizontal. As 197 pyrite has a cubic crystal structure, [100], [010], and [001] axes are equivalent and 198 orthogonal, and the maximum misorientation imposed by symmetry is 62.8°. The pyrite $(m3)^{53}$ match unit (a = 5.4166Å) was used as a reference file for the indexing of the EBSD 199 patterns, the high quality of which was attested by a mean angular deviation (MAD) value 200 201 ranging between 0.2 and 0.6. Six to ten EBSD patterns per grain were collected and the 202 average crystallographic orientations represented in an upper hemisphere equal area pole 203 figure of Euler angle triplets (ϕ_1, ϕ, ϕ_2). For more information on the basics of the EBSD technique, the reader is referred to Prior et al.⁵³. 204

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2.4. SIMS settings:

The samples were stored in the vacuum chamber at 2.5×10^{-8} mbar to 2×10^{-9} mbar. 207 Samples were sputtered with a 3nA ¹⁶O⁻ primary beam accelerated by a 13kV high voltage. 208 209 The corresponding spot size was 2.5µm to 3µm (Figure 2). The primary beam was mainly 210 focused through L3 and L4 lenses of the primary column in Gaussian mode and the 211 aberration was reduced by using a PBMF aperture of 200µm. The L4 aperture was maintained open at 750µm. These settings of primary beam apertures were the same for both 212 213 instruments and the main SIMS settings are summarized in Table 1. The entrance slit was set 214 to 61µm at SwissSIMS and 85µm at CRPG-IPNT, corresponding to a mass resolving power 215 (MRP) of M/ Δ M~ 6700-7000 (slit 3) at SwissSIMS and M/ Δ M~ 6100 (slit 3) at CRPG-IPNT 216 resolving the interference of ${}^{53}CrH^+$ on ${}^{54}Fe^+$ (M/ ΔM = 6088). The interference of ${}^{54}Cr^+$ on 217 ⁵⁴Fe⁺ (MRP~74,000) was monitored by analyzing ⁵²Cr⁺ as described elsewhere⁴⁵. Although pyrite standards have no detectable Cr, hundreds to a thousand of counts of ⁵²Cr are measured 218 219 in natural pyrites, calling for a need to estimate the ⁵⁴Cr⁺ contribution at mass 54 and then 220 correct for it. The correction scheme for Cr was established using a magnetite standard, 221 which has a detectable amount of Cr, resulting in a final correction of ~0.15‰ on δ^{56} Fe per 1×10^4 counts per second (cps) of ${}^{52}Cr^+$. The interference of ${}^{56}FeH^+$ on ${}^{57}Fe^+$, which requires a 222 223 MRP of 7700 could not be clearly resolved with the chosen slit settings. However, this high spatial resolution method is developed in order to analyze δ^{56} Fe values in micrometric 224

pyrites. To obtain a maximum internal error of 0.3‰ (2SE standard error) on δ^{56} Fe, the 225 minimum 56 Fe⁺ intensity on the pyrite standard should be $4x10^7$ cps, which cannot be attained 226 227 under MRP>7000. Pyrite being nominally anhydrous mineral held under the chamber vacuum of $\sim 2x10^{-9}$ mbar yields low interferences of 56 FeH⁺ with the 57 Fe⁺. When pyrites are 228 229 prepared in an indium mount, the iron hydride was not detected on the faraday cup (FC) 230 measurements (see section 2.5). The secondary ion beam was filtered by an energy slit of 50 231 eV. A 2000µm square field aperture was used to clip ~10% of the signal and to remove off-232 axis aberrations of the secondary ion beam. We used a transmission setting (Maximum Area 233 (MA) 80) leading to a field of view of the sample of 20x20µm in the field aperture. The secondary ⁵⁴Fe⁺, ⁵⁶Fe⁺, ⁵⁷Fe⁺ and ⁵²Cr⁺ ions were accelerated at 10kV and analyzed on three 234 235 off-axis FCs and one electron multiplier (EM) (detectors C, H1, H'2 and L2 respectively). 236 The C and H1 FCs were equipped with $10^{11}\Omega$ resistors. The H'2 FC was equipped with a $10^{12}\Omega$ resistor at CRPG-IPNT and a $10^{11}\Omega$ resistor at SwissSIMS. The relative yields of the 237 238 amplifiers of the FCs were calibrated before each session on both ion microprobes and the 239 background noises of the FCs were measured for each analysis. The high voltage of the EM 240 (EM HV) was adjusted before each session. No aging effect of the EM was observed during 241 the session. Presputtering time is necessary to remove the 35nm of gold and potential surface 242 contamination, to implant primary ions in the sample surface and to get a stable secondary signal. The intensity of ⁵⁶Fe⁺ increased until it became stable after 90s (Figure S2, 243 244 supporting information B). The intensity of the signal is then stable with a typical count 245 rate between $4x10^7$ and $5x10^7$ cps when using a 3nA primary beam intensity.

246 The analytical routine then consisted of 90s of presputtering followed by 60 cycles (5s 247 each) of collection separated by 0.08s waiting time, for a total of 7 min per analysis. After 248 presputtering, automatic beam centering in the field and contrast apertures, the energy slit and transfer deflectors were performed routinely. The typical count rate of ⁵⁴Fe⁺, ⁵⁶Fe⁺ and ⁵⁷Fe⁺ 249 250 are 3.2x10⁶, 4.8x10⁷ and 1.1x10⁶cps respectively for Balmat reference material measured at 251 the SwissSIMS instrument tuned with a primary beam intensity of 3.05nA and 4.3x10⁶, 6.4x10⁷ and 1.4x10⁶cps, respectively, at the CRPG-IPNT instrument tuned with a primary 252 253 beam intensity ranging from 3.2 to 3.5nA (Table 1). The difference in ion counting 254 intensities between the two instruments is due to the higher primary current set at CRPG-255 IPNT and also reflects a differential transmission due to the use of a larger entrance slit at CRPG-IPNT. The internal precision on δ^{56} Fe of Balmat standard was $\pm 0.10\%$ (2SE). The 256 257 reproducibility is reported in terms of 2SD, standard deviation. The external reproducibility

258 ranges between $\pm 0.24\%$ (n=33) to $\pm 0.30\%$ (n=17, 2SD) at CRPG-IPNT and from 259 $\pm 0.15\%$ (n=10) to $\pm 0.28\%$ (n=39, 2SD) at SwissSIMS (**Table 2**).

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2.5. *Mass Interferences* (⁵³CrH⁺ and ⁵⁶FeH⁺):

The required MRP to resolve the isobaric interference of ${}^{54}Cr^+$ on ${}^{54}Fe^+$ is out of the 262 ion microprobe capabilities (MRP~74,000), thus, the interference was indirectly quantified 263 by measuring ⁵²Cr⁺. The detailed procedure for Cr correction⁴⁵ shows that the ⁵³Cr/⁵²Cr ratio 264 measured by SIMS is similar to that determined from Cr isotopes natural abundances 265 (${}^{52}Cr=83.8\%$, ${}^{53}Cr=9.5\%$ and ${}^{54}Cr=2.4\%$). We thus used the natural abundances of Cr 266 isotopes combined with the measured ${}^{52}Cr^+$ intensity to calculate the ${}^{54}Cr^+$ intensity in 267 standards and samples and then corrected the ⁵⁴Cr contribution from the measured ⁵⁴Fe 268 signal. The ⁵³CrH⁺ peak height was measured in Russie magnetite standard, on the axial EM 269 270 (monocollection mode), using a 800pA primary beam intensity to obtain $\sim 2x10^5$ cps on 54 Fe⁺ 271 and to not saturate the detector. Under a vacuum of 3.4x10⁻⁹mbar, ⁵³CrH⁺ peak represents less than 0.05% of the ⁵⁴Fe⁺ peak (Figure 1A). Tens of counts are detected for ⁵²Cr⁺ in pyrite 272 273 standards using a 3nA primary beam, meaning that this hydride contribution can be ignored.

Accuracy of analysis can also be impacted by the interference of ⁵⁶FeH⁺ on ⁵⁷Fe⁺. In 274 275 this study, the MRP was set lower (~6800) than that required to separate these two species. Getting a higher MRP to have accurate δ^{57} Fe data would decrease the precision of 56 Fe/ 54 Fe 276 ratios. However, the contribution from ⁵⁶FeH⁺ hydride has been measured to evaluate the 277 reliability of ⁵⁷Fe/⁵⁴Fe ratios. High resolution scan (MRP 7800) of the mass ⁵⁷Fe was carried 278 279 out on the axial EM using a 1nA primary beam intensity (Figure 1C). The magnitude of ⁵⁶FeH⁺ peak is 0.05% of the ⁵⁷Fe⁺ peak height due to good vacuum conditions in the analysis 280 chamber (~3.4x10⁻⁹mbar) but can be up to 0.2% of the ⁵⁷Fe⁺ peak height when vacuum 281 conditions deteriorate to 2x10⁻⁸mbar. The level of ⁵⁶FeH⁺ formation is thus estimated to 282 0.01‰-0.04‰. At MRP 6800, the mass scan shows that ⁵⁷Fe flat top is affected by the tail of 283 the hydride peak (Figure 1B). The hydride contribution on ⁵⁷Fe signal is thus insignificant 284 for measurements done in indium mounts. The contribution of hydrides to the Fe isotopic 285 signal, in particular the interference of ⁵⁶FeH⁺ with ⁵⁷Fe⁺, can also be evaluated by the 286 relationship between the ⁵⁶Fe/⁵⁴Fe and ⁵⁷Fe/⁵⁴Fe ratios. Pyrites from sedimentary rocks 287 288 (Sonoma basin, USA) and standards (Balmat pyrite) measured during the July 2020 session are plotted in a three-isotopes diagram using the natural logarithm of the measured 56 Fe/ 54 Fe 289 and ⁵⁷Fe/⁵⁴Fe ratios (Figure 1D, data available in Table S1, supporting information B). All 290 291 the data define a slope of 0.679 (± 0.007) with a correlation coefficient r² of 0.987, which is

292 consistent with the expected mass-dependent fractionation slope of 0.678. Since the measured 293 slope is consistent with the terrestrial mass fractionation slope, the contribution of ⁵⁶FeH⁺ to 294 the ⁵⁷Fe/⁵⁴Fe ratio is assumed negligible. Sample mounting using indium rather than epoxy 295 together with vacuum conditions below $5x10^{-9}$ mbar are crucial for maintaining small hydride 296 contributions and to produce reliable δ^{57} Fe data and high precision δ^{56} Fe values.

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 - 2.6. IMF correction:
- The effect of a mass dependent fractionation due to the instrument, or instrumentalmass fractionation (IMF), is defined as:

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$$\alpha_{inst} = \frac{\left(\frac{5^{5}Fe}{5^{4}Fe}\right)_{measured}}{\left(\frac{5^{5}Fe}{5^{4}Fe}\right)_{true}}$$
 or $\Delta^{56}Fe$ inst (%) = $\delta^{56}Fe$ measured $-\delta^{56}Fe$ true by using the

302 approximation $\Delta^{56}Fe_{inst} \approx \ln (\alpha_{inst})$.

The "measured" and "true" δ^{56} Fe values are iron isotopic composition measured in the 303 304 same reference material by SIMS and MC-ICP-MS. The typical IMF on pyrite is ~ $-30 \pm 5\%$, 305 measured by both Hyperion-II and Duoplasmatron sources but the exact value depends on the 306 instrument tuning. For example, a 4‰ shift in the IMF for Balmat pyrite was measured after 307 retuning of the primary beam. Slight modifications of the primary beam (i.e. different high 308 voltages on the primary lenses resulting in similar primary intensity but different beam 309 densities) and entrance slit settings lead to the variation of the IMF on Balmat pyrite between 310 -31.20 and -33.10‰ at CRPG-IPNT and highly variable, from -27.23 to -35.16‰ at 311 SwissSIMS (Table 2). We measured the IMF using Balmat pyrite standard to monitor the 312 stability of the ion probe during each session.

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- 314 **3. Results and discussion**
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- 316 *3.1. Spot size:*

The resolution of ion images of a silicon grid on an electron multiplier is used to estimate the ¹⁶O⁻ primary beam size. To find the best possible resolution of the image, (i.e., getting the smallest beam) we tested different combinations of primary lenses, and the best result was obtained by setting the voltage on L1 and L2 to 9800V and 9900V, respectively, and keeping the L3 close to 8500V. The results of the ³⁰Si⁺ ion images are reported in **Figure 2A**. Using a 3nA ¹⁶O⁻ beam, the ability to differentiate two silicon bars on the 3µm grid indicates a spot comparable to 3µm size. The 2µm gap between the two horizontal Si bars is 324 not completely resolved whereas the vertical bars are clearly visible, meaning the spot size is 325 between 2.5 and 3µm. The spot size was verified with a scanning electron microscope (SEM) 326 (Figure 2B) and an interferometric microscope (Figure 2C) after the SIMS measurements of 327 a sedimentary micro-pyrite and the pyrite standard (Balmat). White light interferometric 328 microscopy allows for the precise quantification of the beam size, by imaging the relief of the 329 beam crater in both X and Y directions (Figures 2C and 2D). The primary beam size is 330 measured at the bottom of the pit and corresponds to a $\sim 3\mu m$ diameter spot. This quantitative 331 analysis is consistent with the beam size estimated through ion imaging of the Si grid. The 332 sputtered area is extended over a diameter of 6 x 7-9µm depending on the session and the pit 333 is $\sim 3\mu m$ deep (Figure 2D). The asymmetric shape of the crater and the larger sputtered area 334 compared to the real primary beam size are due to the incidence angle of the beam with the sample surface, which is characteristic of the Cameca[®] ion probes (except NanoSIMS) 335 336 (Figure 2B).

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3.2. Crystal orientation and topography effects on instrumental mass fractionation (IMF):

The IMF is the main limitation in the accuracy of SIMS analysis^{54,55}. It results from 340 various processes occurring during secondary atom ionization, extraction, secondary ion 341 transmission and detection⁵⁵. The IMF is known to be greatly dependent on the sample 342 343 characteristic (mineralogy, chemical composition and crystallographic orientation⁵⁵⁻⁵⁷). This 344 effect is corrected by measuring reference material (same mineral, crystallography and major element chemical compositions) and samples in the same analytical conditions. The IMF 345 variations occur in various isotopic systems, for example, the $\delta^{18}O$ measurements of garnets 346 are strongly affected by their Ca-Fe-Mg content^{58,59}, as well as δ^{34} S in Fe-Ni sulfides^{56,60,61}, 347 and Mg and Si isotopes in silicates (e.g. olivine, glass, pyroxene) with an IMF depending on 348 their Mg content^{43,62}. Crystal orientation may also have in some cases a strong influence on 349 350 the IMF, as demonstrated for example on Fe isotopic compositions in magnetite⁵⁷, on S 351 isotope compositions in sphalerite and galena⁶³ and on U-Pb dating in baddeleyite⁴.

As pyrites are not affected by major element substitution, *i.e.* no solid solution or chemical variability, the potential variations of the IMF can only be the result of the crystal orientation and/or the topography. The EBSD pole figure shown in **Figure 3B** displays the crystallographic orientations of the three different grains of Balmat standard projected on a plane (XY) parallel to the surfaces analyzed by SIMS (and EBSD). With misorientation angles between [100] axes in Grain 1, 2, and 3 (G1-G3) and the reference direction Z (i.e. the 358 normal to the page pointing towards the reader) of 14°, 6°, and 1°, respectively, the analyzed 359 surfaces can be considered nearly parallel to the face of the pyrite unit cell (i.e. normal to the 360 [100] axis). The misorientation angle between the [100] axes located in the middle of the pole figure is 18° between G1 and G2, and 15° between G1 and G3. Fe isotopic measurements 361 362 show respective mean δ^{56} Fe values and external reproducibility of -0.29 ±0.30‰ (2SD) / 363 $\pm 0.13\%$ (2SE, n=5), -0.59 $\pm 0.42\%$ (2SD) / $\pm 0.19\%$ (2SE, n=5) and -0.32 $\pm 0.44\%$ (2SD) / 364 $\pm 0.20\%$ (2SE, n=5), for G1, G2, and G3 (Figure 3A). Those δ^{56} Fe values suggest that the inter-grain variability is lower than the reproducibility (2SD) and the uncertainty on the 365 366 averages (2SE). Even though the EBSD measurements on a sample set of only three grains have no statistical significance, it can be said at this stage that no obvious relationship 367 368 between the crystallographic orientation of pyrite and SIMS δ^{56} Fe measurements has been observed. A thorough review (beyond the scope of this study) based on a wide range of 369 370 crystallographic orientations is needed to confirm this initial statement. Our results are also 371 consistent with the absence of crystal orientation effects on S isotopes^{57,65}.

372 The channeling effect of the primary ion beam as a function of atomic planes 373 orientation has been shown to influence the secondary ion yields and thus the instrumental mass fractionation^{66,67}. Similarly to magnetite that exhibits channeling effects and plane-374 375 specific IMF for Fe and O isotopes⁵⁷, pyrites are cubic minerals that could experience similar 376 effects. This effect was evaluated by rotating the mount in the sample holder by 90° , 180° 377 and 270°. We ran 3 to 6 analyses per rotation (Table S2, supporting information B). The mean IMF-corrected δ^{56} Feirmmold values for P0, P1, P2 and P3 are -0.38 ±0.31‰ (2SD) / 378 379 ±0.13‰ (2SE, n=6) for P0, -0.49 ±0.37‰ (2SD) / ±0.21‰ (2SE, n=3) for P1, -0.38 ±0.12‰ $(2SD) / \pm 0.07\%$ (2SE, n=3) for P2 and -0.39 $\pm 0.36\%$ (2SD) / $\pm 0.16\%$ (2SE, n=5) (Figure 380 381 4). Considering the external reproducibility (2SD) and the internal error (2SE) together, those 382 data show a similar mean δ^{56} Fe values across the four positions in the holder.

383 Surface topography could also induce artificial iron isotopic variations, especially 384 when pyrite grains are just slightly bigger than the primary spot size. Here, a core to rim profile on Balmat pyrites was performed in order to examine the edge effect on the reliability 385 of δ^{56} Fe analyses. Fe isotope analyses show similar δ^{56} Fe values between the core and the rim 386 of δ^{56} FeBalmatPf1@01= -0.27 ±0.11‰ (2SE, n=1, core) and δ^{56} FeBalmatPf1@8= -0.28 ±0.12‰ (2SE, 387 388 n=1, rim) and an external reproducibility of $\pm 0.18\%$ (2SD) (Figure 5D, data available in 389 Table S2 supporting information B). This profile (Pf1) is characterized by a topographic 390 difference of 1.7µm (Figure 5C), which is not significant to introduce a bias. However, the 391 last analysis is located at $\sim 20 \mu m$ from the grain edge that is $\sim 7 \mu m$ above the enclosing

indium. Thus, this value was measured in the slightly tilted shade zone on the edge of the pyrite (**Figure 5A and Figure 5B**), demonstrating the reliability of δ^{56} Fe values.

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- *395 3.3. Sensitivity*:

The sensitivity depends on the sputtering time, ionization, extraction of the Fe⁺ ion from the matrix and secondary ions transmission until the detectors. It is defined as count rate per ppm of Fe in the analyzed phase per nA of the primary beam (cps/ppm/nA). As the Fe content is constant in pyrites, the expression of the sensitivity is approximately proportional to the ion yield:

401 56 Fe yield = 56 Fe⁺ / P intensity.

402 The sensitivity calculation is commonly used to evaluate the transmission of an ion microprobe. **Table 1** shows the ⁵⁶⁺Fe ion yields on Balmat pyrite, obtained over the different 403 sessions at CRPG-IPNT and SwissSIMS equipped with the Hyperion-II source. We 404 405 compared these results to the sensitivity obtained with the conventional Duoplasmatron source^{45,46}. The sensitivities determined on Balmat pyrite range from 1.56x10⁷ to 406 2.01x107cps/nA at CRPG-IPNT. Sessions performed at SwissSIMS (July 2019, January 407 408 2020, March 2020 and September 2020) show similar sensitivities, ranging from 409 1.49x10⁷cps/nA to 1.61x10⁷cps/nA. Higher sensitivities obtained at CRPG-IPNT compared 410 to SwissSIMS are explained by different widths of the entrance slit. Sensitivities obtained with the Duoplasmatron source vary from 1.2x10⁷cps/nA⁴⁵ to 1.5x10⁷cps/nA⁴⁶. However, the 411 resulting transmissions in the two Duoplasmatron-based studies are not directly comparable 412 413 as the width of the field aperture (FA), the entrance and exit slit and the Maximum Area 414 (MA) are different between these two studies (Table 1). In order to compare sensitivities 415 obtained by Hyperion and Duoplasmatron, we performed two tests using (1) a MA 80 and field aperture closed at ~2500 μ m (comparable with sensitivity obtained previously⁴⁵) and (2) 416 417 a MA 160 and an opened field aperture to transmit 100% of the signal⁴⁶. Using a MA of 80 and similar FA, entrance and exit slit widths, the ⁵⁶Fe sensitivity is 1.56x10⁷cps/nA and better 418 419 than the $1.2 \times 10^7 \text{cps/nA}$ obtained with the Duoplasmatron⁴⁵. The higher sensitivity with the 420 Hyperion is due to the use of a smaller beam, which is less clipped in the field aperture compared to the Duoplasmatron beam. Using a MA 160, we obtained a sensitivity of 421 422 5.05×10^{6} cps/nA, which is lower than the 1.5×10^{7} cps/nA achieved by the Duoplasmatron⁴⁶. 423 However, these sensitivities are not directly comparable as O₂⁻ and O⁻primary beams were respectively used in Whitehouse and Fedo⁴⁶ and in the present study. The higher sensitivity 424 obtained previously⁴⁶ can be thus attributed to the more efficient sputtering rate of the O_2^{-1} 425

426 beam. The use of smaller primary beam currents reduces the size of the crossover and off-427 axis aberrations of the secondary ion beam, which helps to define a sharper slit image. 428 Therefore, for a given MRP, the entrance slit can be more opened using the Hyperion and this 429 results in a gain of sensitivity compared to the Duoplasmatron. This test illustrates the ability 430 of the source to provide enough secondary ion signals with a 3nA beam focused on a 3µm 431 spot and to achieve a better sensitivity than that delivered by Duoplasmatron. The Fe 432 secondary ion signals produced by the 3nA primary beam can be detected by MCFC and thus, provides high precision δ^{56} Fe measurements along with higher spatial resolution. 433

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3.4. Reproducibility and accuracy:

The reproducibility of the δ^{56} Fe measurements on the Balmat reference material was 436 437 established over three sessions (February 2018, April 2018 and September 2020) at CRPG-438 IPNT and four sessions (July 2019, January 2020, March 2020 and June 2020) at SwissSIMS 439 Lausanne (Table 2). Balmat pyrite (same grain) displays a respective long-term 440 reproducibility of ±0.25‰ (2SD) for 166 measurements (Figure 6) at CRPG-IPNT and 441 ±0.22‰ (2SD) for 185 measurements (June 2020) at SwissSIMS ion probe. The short-term 442 reproducibility on Balmat pyrite varied from ±0.24‰ (2SD, February 2018, n=33, Table S3 443 **supporting information B**) to ±0.26‰ (2SD, April 2018, n=133) at CRPG-IPNT and from 444 ±0.18‰ (2SD, March 2020, n=33) to ±0.35‰ (2SD, January 2020, n=16) at SwissSIMS. The 445 reproducibility obtained on SpainCR standard (July 2018) is close to those measured in 446 Balmat pyrite with a value of $\pm 0.28\%$ (2SD, n=61). Published data obtained using the ¹⁶O⁻ Duoplasmatron source⁶⁸ on three days of analysis show a reproducibility of $\pm 0.44\%$ (2SD, 447 448 n=17) on the same grain of Balmat, which highlight the better stability of the Hyperion-II 449 source compared to the Duoplasmatron.

The accuracy of the SIMS technique was tested on two pyrite standards which have different isotopic compositions. We used Balmat as a reference standard and considered SpainCR as an unknown pyrite. The δ^{56} Fe value for SpainCR was determined using the IMF calculated on Balmat reference material. The δ^{56} Fe value for SpainCR pyrite calculated at +0.64 ±0.26‰ (2SD, n=2) is in quite good agreement with the value of δ^{56} Fe=+0.52 ±0.03‰ determined by MC-ICP-MS, demonstrating the accuracy of the SIMS method.

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459 **4.** Conclusions

An ion microprobe equipped with the new Hyperion-II Radio Frequency source is 460 461 able to determine iron isotope ratios with high accuracy, at high precision ($\sim 0.25\%$, 2SD) 462 and high spatial resolution (3µm). We have detailed a procedure to achieve δ^{56} Fe 463 measurements a primary intensity 3 times lower than that traditionally delivered by the Duoplasmatron source, yet we achieved better precision. The MRP was intentionally set at a 464 465 lower value than that required to resolve ⁵⁶FeH⁺ from ⁵⁷Fe⁺ to attain a minimum ⁵⁶Fe⁺ count rate of 4×10^7 cps on a pyrite reference material to produce high precision δ^{56} Fe values. The 466 level of ⁵⁶FeH⁺ is low compared to the intensity of ⁵⁷Fe⁺ under high vacuum. A MRP of 467 ~6700 is sufficient to limit its contribution to 57 Fe signal. In our sample set, we demonstrated 468 the absence of topography and crystal orientation effects. Currently, this new procedure is 469 470 applied to major iron-bearing minerals to better constrain natural iron isotopic variabilities at 471 micrometric scale in sedimentary pyrites. The future investigation of *in situ* δ^{56} Fe signatures 472 in minor iron-bearing phases such as oxides, carbonates and silicates will serve as a 473 promising tool to answer fundamental questions in extraterrestrial and terrestrial petrology 474 and to gain a better understanding of the biogeochemical iron cycles.

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TABLE 1 SIMS settings for each Fe isotopes session at CRPG-IPNT (February 2018,
April 2018 and September 2020) and SwissSIMS (July 2019, January 2020, March 2020 and
June 2020) using the Hyperion-II source and resulting ⁵⁶Fe⁺ intensities and ion yields
measured on Balmat pyrite. Comparison with settings and ⁵⁶Fe⁺ yields measured with the
Duoplasmatron^{45,46}.

					N 7			PUBLI	SHED
			11	HIS STUL	JΥ			DAT	ГА
	C	DDC IDN	T					CRPG-	Nord
	(LKFU-IFN	1		SW188	511/15		IPNT	SIMS
]	Hyperion-l	Π		Нуре	rion-II		Duoplas	matron
Session	Feb.18	Ap.18	Sep.20	Jul.19	Jan.20	Mar.20	Jun.20	Ref. 45	Ref.46
Field Ap.	20	000	2400		20	000		2500	3000
(µm)	20	.00	2400		20	.00		2500	5000
Contrast		400			4	00		40	ົງ
Ap. (μm)		100				00			
Max. Area	8	30	80		8	30		80	160
			160						100
Entrance	8	35	61	61				-	60
Slit (µm)			01						
Energy slit		50		50				50	60
(eV)					-	-			
Slit Mode		Circular		Circular				Circu	ılar
Exit slit	1	50um (slit	3)		150um	(slit 3)		150µm	250µm
	-	copini (bito			10 0 μ	(511 0)		(slit 3)	(slit 2)
MRP	61	.00	6800	6700-6800				~7000	<6000
Primary									
Intensity	3.40	3.70	3.00	3.14	3.05	2.99	2.80	10	10
(nA)									
⁵⁶ Fe ⁺ count									
rate on	6.5E+7	7.45E+7	4.70E+7	4.90E+7	4.85E+7	4.46E+7	4.49E+7	~ 1.2E+8	~1.5E+8
Balmat			1.51E+7						
(cps)									
⁵⁶ Fe ⁺ yield			1.56E+7						
on Balmat	1.87E+7	2.01E+7	5.05E+6	1.56E+7	1.59E+7	1.49E+7	1.61E+7	~1.2E+7	1.5E+7
(cps/nA)									

TABLE 2 True δ^{56} Fe values of the standards (Balmat and SpainCR pyrites) and corresponding IMF during the different sessions at CRPG-IPNT and SwissSIMS and comparison with published IMF obtained with the Duoplasmatron⁴⁵. True δ^{56} Fe value of SpainCR standard was determined by MC-ICP-MS method at CRPG using a reported procedure⁵⁰. The reproducibility is reported as 2SD, standard deviation. n= number of analysis; *n.a*= not available data.

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INSTRUM	IENTAL	MASS FI	RACTIO	NATION	δ ⁵⁶ Fe _{IMF}	AND RE	PRODU	CIBILITY	(2SD, ‰)
				(#ANA	LYSIS)				
	CF	RPG-IPNT				Swis	sSIMS		CRPG-
	H	yperion-II				Нуре	rion-II		IPNT
									Duoplas
									matron
Session	Feb.	Apr.	Ju.	Sept.	Ju.	Jan.	Mar.	Jun.	ref.45
56551011	2018	2018	2018	2020	2019	2020	2020	2020	
Ralmat	-33.10	-32.47		-31.20	-35.16	-27.23	-32.34	-33.06	-32.00
Dannat	±0.24	±0.26	n.a	±0.30	±0.28	±0.35	±0.18	±0.22	±0.20
ry.	(n=33)	(n=133)		(n=17)	(n=41)	(n=16)	(n=33)	(n=185)	
SnainCR			-33.21		-35.50				
Pv	n.a	n.a	±0.28	n.a	±0.32	n.a	n.a	n.a	n.a
1 J.			(n=68)		(n=2)				



FIGURE 1 A) Scan of the ⁵⁴Fe signal on the axial EM, using a mass resolution of 6800 to reveal ⁵³CrH⁺ contribution. This scan was made on Russie magnetite standard which have a detectable Cr content. B) Scan of the ⁵⁷Fe signal on the axial EM, using a mass resolution of 6800 to reveal ⁵⁶FeH⁺ contribution. C) High mass resolution (MRP 7800) scan of the ⁵⁷Fe signal, where ⁵⁷Fe⁺ and ⁵⁶FeH⁺ peaks are separated. D) Three-isotope plot of the logarithm of the measured Fe isotope ratios in Balmat pyrite standard (white dots) and pyrites from sedimentary rocks (blue dots), corrected from the ⁵⁴Cr contribution (July 2020 session at SwissSIMS). The regression line gives a slope of 0.679 ± 0.007 , which is in good agreement with the theoretical slope of 0.678 within the error bar. The theoretical value represents the mass dependent fractionation law for Fe isotopes using a simple harmonic oscillator approximation. Data are available in Table S1 (supporting information B).



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A) ³⁰Si⁺ image of the Si-grid on electron multiplier. The spatial resolution of 780 FIGURE 2 the primary ¹⁶O⁻ beam is determined according to the ability to differentiate two silicon bars 781 on the 4µm, 3µm or 2µm grid. B) SEM image of a sedimentary micro-pyrite after SIMS 782 783 analyses. The shape of the spot is due to the position of the source which makes an angle with 784 the sample surface. C) Image from interferometric microscope of the sputtering pit. D) X and 785 Y topographic profiles measured by interferometric microscope. The profiles refer to the 786 Figure 2C and show the diameter of the pit, which corresponds to the real spot size (measured 787 at the bottom of the pit), and the sputtered area (blue area on Figure 2D and dashed red circle 788 on Figure 2C, measured at the top of the pit).



FIGURE 3 Crystal orientation (September 2020) tests in Balmat pyrite standard. A) δ^{56} Fe corrected from instrumental fractionation measured in three grains prepared in an indium mount. The reproducibility is given at 2SD and show no clear difference between the grains. Data are available in **Table S2** (**supporting information**). B) Upper hemisphere equal area (i.e. with Z pointing to the reader) EBSD pole figure showing the averaged crystallographic orientations of {100} for the three pyrite grains of interest. Note the strong clustering of [100] axes close to Z.

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FIGURE 4 δ^{56} Fe values IMF-corrected of Balmat pyrite standard rotated by 90° (P1), 180° (P2) and 270° (P3) compared to the initial position (P0). The external reproducibility is ±0.30‰ (2SD) and the internal variability are ±0.10‰ to ±0.20‰ (2SE), allowing to rule out orientation (channeling) effect on IMF variations. Data are available in **Table S2** (supporting information).

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812 Profiles in Balmat reference material. A) Interferometric microscope image FIGURE 5 showing Pf1 profile in the pyrite grain. B) Interferometric microscope 3D image of the pyrite 813 814 standard. C) Topographic profile following Pf1 transect. This profile shows a topographic 815 difference of ~1.5µm between the core (red star) and the rim (green star) of the grain. The 816 gap after the green star (\sim 7µm) highlights the boundary between the pyrite grain and the 817 indium. D) δ^{56} Fe values IMF-corrected in a core to rim profile performed in Balmat pyrite standard. Dashed black line is the true δ^{56} Fe value of Balmat standard (δ^{56} Fe= -0.399‰). The 818 819 uncertainty on average is $\pm 0.09\%$ (2SE) and the external reproducibility is $\pm 0.18\%$ (2SD, 820 grey area). Data are available in **Table S2** (supporting information).

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FIGURE 6 Long term reproducibility on Balmat pyrite reference material (0.25‰, 2SD). Dots are δ^{56} Fe measured with Hyperion-II Radio-Frequency plasma source in February 2018 (blue dots) and April 2018 (white dots) session. Grey diamonds are δ^{56} Fe data from the Duoplasmatron source⁶⁸. Dash black line indicates the true δ^{56} Fe value for Balmat (δ^{56} Fe -(0.399%) and grey area represents the long-term reproducibility of ± 0.25 at 2SD, standard deviation. February 2018 data are available in Table S3 (supporting information B).

845 **Supporting information B:** 846 847 **Electron Probe Microanalysis (EPMA) settings:** 848 The acceleration voltage was 15kV and beam current was 15nA focused in 3µm. 849 Reference materials, including sulfides, oxides and silicates, were tested before the analysis 850 of S, Fe, Co, Mn, Pb, Ti, Cr, Zn, Cu and Ni. Detection limits were 133ppm for S, 130ppm for Fe, 141ppm for Co, 129ppm for Mn, 327ppm for Pb, 79ppm for Ti, 145ppm for Cr, 238ppm 851 852 for Zn, 199ppm for Cu and 150ppm for Ni.

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854 **Supplementary figures:**





857 Reproducibility on Spain standard (n=68) during the July 2018 session 858 FIGURE S1 (CRPG-IPNT) in three different grains. The reproducibility is ±0.28‰ (2SD) and was 859 860 obtained by bracketing method. The black dashed line represents the true value of the 861 standard (δ^{56} Fe=+0.52‰).



864 FIGURE S2 Intensity of 56 Fe⁺ in Balmat pyrite reference material obtained with Hyperion 865 RF using a 3nA primary beam. The 56 Fe⁺ intensity increases and stabilizes after 90s, which 866 corresponds to the necessary presputtering time before the analyses.

A nolucia Nome	$\delta^{56}Fe_{corr}$	1SE	$\delta^{57}Fe_{corr}$	JCE	⁵⁶ Fe ⁺ intensity	ln(^{56/54} Fe	ln(^{57/54} Fe
Analysis Maine	54 _{Cr}	25E	⁵⁴ Cr	25E	(cps)	meas)	meas)
Standards							
Balmat@8	-33.68	0.14	-50.17	0.36	3.91E+07	2.7199	-1.0647
Balmat@9	-34.10	0.13	-50.44	0.40	3.95E+07	2.7195	-1.0650
Balmat@10	-34.09	0.15	-50.63	0.41	3.95E+07	2.7195	-1.0652
Balmat@11	-33.90	0.16	-50.17	0.40	3.91E+07	2.7197	-1.0647
Balmat@12	-33.47	0.12	-49.49	0.37	4.12E+07	2.7201	-1.0640
Balmat@13	-33.69	0.12	-49.87	0.37	4.06E+07	2.7199	-1.0644
Balmat@14	-33.74	0.14	-49.77	0.39	4.08E+07	2.7198	-1.0643
Balmat@15	-33.75	0.14	-49.72	0.33	4.05E+07	2.7198	-1.0643
Balmat@16	-33.62	0.13	-49.55	0.34	4.02E+07	2.7199	-1.0641
Balmat@17	-34.10	0.13	-51.09	0.41	4.53E+07	2.7195	-1.0656
Balmat@18	-34.09	0.11	-50.94	0.30	4.54E+07	2.7195	-1.0655
Balmat@19	-34.16	0.14	-51.15	0.32	4.54E+07	2.7194	-1.0657
Balmat@20	-33.98	0.13	-50.89	0.33	4.47E+07	2.7196	-1.0654
Balmat@21	-34.16	0.13	-50.91	0.29	4.47E+07	2.7194	-1.0654
BalmatMI83@01	-33.61	0.11	-49.93	0.38	4.09E+07	2.7200	-1.0645
BalmatMI83@10	-34.12	0.13	-50.40	0.36	4.29E+07	2.7194	-1.0649
BalmatMI83@11	-34.14	0.12	-50.32	0.37	4.28E+07	2.7194	-1.0649
BalmatMI83@12	-34.06	0.15	-50.37	0.39	4.36E+07	2.7195	-1.0649
BalmatMI83@13	-33.76	0.10	-50.07	0.45	4.29E+07	2.7198	-1.0646
BalmatMI83@14	-34.14	0.12	-50.60	0.32	4.33E+07	2.7194	-1.0651
BalmatMI83@15	-34.00	0.12	-50.56	0.37	4.23E+07	2.7196	-1.0651
BalmatMI83@16	-33.79	0.12	-50.26	0.34	4.37E+07	2.7198	-1.0648
BalmatMI83@17	-34.12	0.12	-51.02	0.38	4.32E+07	2.7194	-1.0656
BalmatMI83@18	-33.77	0.13	-50.34	0.37	4.36E+07	2.7198	-1.0649
BalmatMI83@19	-33.90	0.11	-50.22	0.31	4.31E+07	2.7197	-1.0648
BalmatMI83@2	-33.45	0.13	-49.54	0.36	4.12E+07	2.7201	-1.0641
BalmatMI83@3	-33.68	0.12	-49.97	0.36	4.03E+07	2.7199	-1.0645
BalmatMI83@4	-33.86	0.11	-50.18	0.33	4.08E+07	2.7197	-1.0647
BalmatMI83@5	-33.75	0.11	-50.03	0.38	4.07E+07	2.7198	-1.0646
BalmatMI83@6	-33.72	0.11	-50.00	0.40	4.09E+07	2.7198	-1.0645

BalmatMI83@7	-34.13	0.12	-50.28	0.30	4.52E+07	2.7194	-1.0648
BalmatSTWanas@01	-33.83	0.12	-50.04	0.32	4.31E+07	2.7197	-1.0646
BalmatSTWanas@10	-33.98	0.11	-50.75	0.37	4.28E+07	2.7196	-1.0653
BalmatSTWanas@11	-33.99	0.11	-50.85	0.45	4.28E+07	2.7196	-1.0654
BalmatSTWanas@12	-33.56	0.13	-50.12	0.42	4.28E+07	2.7200	-1.0647
BalmatSTWanas@13	-33.77	0.11	-50.04	0.37	4.31E+07	2.7198	-1.0646
BalmatSTWanas@15	-33.69	0.14	-49.95	0.37	4.29E+07	2.7199	-1.0645
BalmatSTWanas@16	-33.73	0.12	-50.13	0.36	4.28E+07	2.7198	-1.0647
BalmatSTWanas@17	-33.94	0.14	-50.65	0.42	4.21E+07	2.7196	-1.0652
BalmatSTWanas@2	-34.04	0.14	-50.66	0.36	4.26E+07	2.7195	-1.0652
BalmatSTWanas@3	-34.23	0.11	-51.01	0.33	4.25E+07	2.7193	-1.0655
BalmatSTWanas@4	-34.15	0.12	-50.90	0.34	4.27E+07	2.7194	-1.0654
BalmatSTWanas@5	-33.90	0.13	-50.14	0.35	4.28E+07	2.7197	-1.0647
BalmatSTWanas@6	-34.01	0.10	-50.39	0.41	4.25E+07	2.7196	-1.0649
BalmatSTWanas@7	-33.85	0.11	-50.29	0.39	4.28E+07	2.7197	-1.0648
BalmatSTWanas@8	-33.96	0.10	-50.48	0.39	4.30E+07	2.7196	-1.0650
BalmatSTWanas@9	-33.77	0.11	-50.33	0.36	4.34E+07	2.7198	-1.0649
Samples							
MI83@01	-32.96	0.14	-49.19	0.41	3.76E+07	2.7206	-1.0637
MI83@10	-32.41	0.17	-48.05	0.37	3.74E+07	2.7211	-1.0626
MI83@11	-32.54	0.13	-48.14	0.35	3.88E+07	2.7210	-1.0627
MI83@12	-33.13	0.11	-48.83	0.36	3.83E+07	2.7204	-1.0634
MI83@13	-32.33	0.15	-47.79	0.39	3.67E+07	2.7212	-1.0623
MI83@14	-33.10	0.15	-49.01	0.44	3.73E+07	2.7205	-1.0635
MI83@15	-31.74	0.13	-46.88	0.40	3.56E+07	2.7218	-1.0614
MI83@16	-32.42	0.11	-47.95	0.39	3.89E+07	2.7211	-1.0625
MI83@17	-33.12	0.13	-48.78	0.42	3.84E+07	2.7204	-1.0633
MI83@18	-32.08	0.16	-47.28	0.44	3.80E+07	2.7215	-1.0618
MI83@19	-32.73	0.12	-48.35	0.37	3.90E+07	2.7208	-1.0629
MI83@2	-32.90	0.11	-48.63	0.34	3.83E+07	2.7207	-1.0632
MI83@20	-32.54	0.15	-48.32	0.44	3.78E+07	2.7210	-1.0629
MI83@21	-31.55	0.14	-46.73	0.37	3.80E+07	2.7220	-1.0613
MI83@22	-32.32	0.11	-47.94	0.35	4.15E+07	2.7212	-1.0625

MI83@23	-31.56	0.20	-47.04	0.51	3.82E+07	2.7220	-1.0616
MI83@24	-32.09	0.15	-47.80	0.42	4.03E+07	2.7215	-1.0623
MI83@25	-31.84	0.12	-47.15	0.36	4.11E+07	2.7217	-1.0617
MI83@26	-32.03	0.13	-47.58	0.37	4.04E+07	2.7215	-1.0621
MI83@27	-31.26	0.14	-46.82	0.39	4.13E+07	2.7223	-1.0614
MI83@28	-31.59	0.13	-46.97	0.33	4.14E+07	2.7220	-1.0615
MI83@29	-32.34	0.13	-48.15	0.30	4.20E+07	2.7212	-1.0627
MI83@3	-32.00	0.13	-47.57	0.34	3.94E+07	2.7216	-1.0621
MI83@30	-31.68	0.14	-47.34	0.42	4.11E+07	2.7219	-1.0619
MI83@31	-32.61	0.13	-48.28	0.33	4.18E+07	2.7209	-1.0628
MI83@32	-32.20	0.15	-47.84	0.41	3.82E+07	2.7214	-1.0624
MI83@33	-32.65	0.19	-48.18	0.44	3.23E+07	2.7209	-1.0627
MI83@34	-32.66	0.12	-48.60	0.35	4.08E+07	2.7209	-1.0631
MI83@35	-32.57	0.16	-48.13	0.39	3.95E+07	2.7210	-1.0627
MI83@36	-32.33	0.14	-47.94	0.32	4.04E+07	2.7212	-1.0625
MI83@37	-32.04	0.12	-47.81	0.39	3.82E+07	2.7215	-1.0624
MI83@39	-31.35	0.12	-46.70	0.35	3.98E+07	2.7222	-1.0612
MI83@4	-32.71	0.15	-48.55	0.40	3.74E+07	2.7208	-1.0631
MI83@40	-30.82	0.12	-45.85	0.36	3.95E+07	2.7227	-1.0604
MI83@41	-29.48	0.34	-43.79	0.56	4.03E+07	2.7240	-1.0583
MI83@42	-31.36	0.14	-47.11	0.40	4.05E+07	2.7222	-1.0616
MI83@43	-31.87	0.11	-47.73	0.40	4.13E+07	2.7217	-1.0623
MI83@44	-30.99	0.19	-45.96	0.51	3.50E+07	2.7225	-1.0605
MI83@7	-32.34	0.14	-47.96	0.38	3.89E+07	2.7212	-1.0625
MI83@8	-32.42	0.12	-47.98	0.34	3.80E+07	2.7211	-1.0625
MI83@9	-32.28	0.13	-47.82	0.42	3.89E+07	2.7213	-1.0624
STWanas@01	-31.35	0.13	-46.69	0.34	4.23E+07	2.7222	-1.0612
STWanas@10	-31.53	0.12	-46.68	0.48	3.81E+07	2.7220	-1.0612
STWanas@11	-30.79	0.13	-45.85	0.38	4.19E+07	2.7228	-1.0604
STWanas@12	-31.11	0.13	-46.67	0.35	4.19E+07	2.7225	-1.0612
STWanas@13	-30.97	0.12	-46.31	0.34	4.22E+07	2.7226	-1.0608
STWanas@14	-31.21	0.12	-46.54	0.37	4.19E+07	2.7223	-1.0611
STWanas@15	-31.33	0.13	-46.61	0.38	4.20E+07	2.7222	-1.0611

STWanas@16	-31.29	0.13	-46.13	0.31	4.20E+07	2.7223	-1.0607
STWanas@17	-31.02	0.14	-46.16	0.38	4.26E+07	2.7225	-1.0607
STWanas@18	-30.36	0.12	-45.26	0.32	4.15E+07	2.7232	-1.0598
STWanas@19	-31.08	0.15	-46.14	0.30	4.15E+07	2.7225	-1.0607
STWanas@2	-30.91	0.11	-45.89	0.32	4.18E+07	2.7227	-1.0604
STWanas@21	-31.17	0.12	-46.37	0.35	4.23E+07	2.7224	-1.0609
STWanas@22	-31.30	0.15	-46.22	0.34	4.19E+07	2.7223	-1.0608
STWanas@23	-31.08	0.13	-46.31	0.30	4.26E+07	2.7225	-1.0608
STWanas@25	-31.21	0.12	-46.55	0.35	4.28E+07	2.7224	-1.0611
STWanas@26	-29.54	0.18	-43.80	0.42	3.64E+07	2.7240	-1.0583
STWanas@29	-31.11	0.13	-46.40	0.32	4.27E+07	2.7225	-1.0609
STWanas@30	-30.35	0.17	-45.14	0.49	3.86E+07	2.7232	-1.0597
STWanas@32	-31.03	0.10	-46.35	0.40	4.21E+07	2.7225	-1.0609
STWanas@33	-31.14	0.13	-46.30	0.40	3.63E+07	2.7224	-1.0608
STWanas@34	-31.56	0.13	-46.95	0.38	4.24E+07	2.7220	-1.0615
STWanas@35	-31.24	0.12	-46.34	0.42	4.23E+07	2.7223	-1.0609
STWanas@36	-31.24	0.14	-46.52	0.40	3.95E+07	2.7223	-1.0611
STWanas@37	-31.18	0.14	-46.42	0.38	4.21E+07	2.7224	-1.0610
STWanas@38	-31.12	0.13	-46.55	0.35	4.22E+07	2.7224	-1.0611
STWanas@39	-30.87	0.13	-46.06	0.37	4.20E+07	2.7227	-1.0606
STWanas@4	-31.62	0.13	-46.99	0.39	4.28E+07	2.7219	-1.0615
STWanas@40	-31.18	0.13	-46.68	0.38	3.92E+07	2.7224	-1.0612
STWanas@5	-31.73	0.13	-47.32	0.29	4.22E+07	2.7218	-1.0619
STWanas@6	-31.44	0.14	-46.86	0.42	4.24E+07	2.7221	-1.0614
STWanas@8	-31.33	0.12	-47.05	0.40	4.23E+07	2.7222	-1.0616
STWanas@9	-30.81	0.12	-45.78	0.32	4.25E+07	2.7228	-1.0603

TABLE S1 δ^{56} Fe and δ^{57} Fe values (corrected from 54 Cr interference), intensity of the signal and logarithm of the measured 56 Fe/ 54 Fe and 57 Fe/ 54 Fe ratios in Balmat pyrite standard and geological samples (pyrites) during July 2020 session (SwissSIMS). The 2SE (standard error) reports the internal uncertainty.

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Analysis Name	δ ⁵⁶ Feraw (‰)	δ ⁵⁶ Fe _{IRMM014} (‰)	2SE
	X-Y te	est	
Position 0° (P0)			
Balmat@25	-31.71	-0.52	0.09
Balmat@26	-31.77	-0.58	0.11
Balmat@27	-31.60	-0.41	0.09
Balmat@31	-31.38	-0.20	0.10
Balmat@32	-31.48	-0.29	0.11
Balmat@33	-31.44	-0.26	0.12
Position 90° (P1)			
Balmat@34	-31.48	-0.30	0.10
Balmat@35	-31.69	-0.50	0.10
Balmat@36	-31.86	-0.67	0.12
Position 180° (P2)			
Balmat@38	-31.49	-0.31	0.10
Balmat@39	-31.60	-0.41	0.10
Balmat@40	-31.60	-0.41	0.12
Position 270° (P3)			
Balmat@45	-31.82	-0.64	0.11
Balmat@46	-31.70	-0.51	0.10
Balmat@47	-31.41	-0.22	0.12
Balmat@48	-31.45	-0.26	0.12
Balmat@49	-31.49	-0.30	0.09
	Crystal orient	ation test	

Grain #1 (G1)			
BalmatG1@38	-29.81	-0.12	0.12
BalmatG1@39	-29.92	-0.23	0.12
BalmatG1@40	-30.21	-0.52	0.10
BalmatG1@41	-29.94	-0.25	0.13
BalmatG1@43	-30.01	-0.32	0.11

Grain #2 (G2)			
BalmatG2@9	-30.25	-0.57	0.11
BalmatG2@10	-30.31	-0.62	0.10
BalmatG2@11	-30.43	-0.74	0.10
BalmatG2@12	-29.93	-0.24	0.10
BalmatG2@13	-30.45	-0.76	0.13
Grain #3 (G3)			
BalmatG3@24	-30.09	-0.41	0.09
BalmatG3@25	-30.20	-0.51	0.10
BalmatG3@26	-30.18	-0.49	0.11
BalmatG3@27	-29.69	0.00	0.10
BalmatG3@28	-29.88	-0.20	0.10
	Profile P	f1	
BalmatPf1@01	-27.37	-0.27	0.11
BalmatPf1@02	-27.62	-0.52	0.12
BalmatPf1@03	-27.52	-0.42	0.12
BalmatPf1@04	-27.60	-0.50	0.10
BalmatPf1@05	-27.50	-0.41	0.11
BalmatPf1@06	-27.49	-0.40	0.13
BalmatPf1@07	-27.49	-0.39	0.12
BalmatPf1@08	-27.38	-0.28	0.12

TABLE S2 Raw δ^{56} Fe and IMF-corrected δ^{56} Fe values measured by SIMS on Balmat pyrite in four different position (0°, 90°, 180° and 270°) to test orientation (channeling) effect (data acquired during September 2020 session); in three Balmat grains mounted in random orientations to test a possible crystal orientation effect (data acquired during September 2020 session); and a core to rim profile in a Balmat grain. The internal uncertainties of the measurements are reported as 2SE.

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Analysis Name	⁵⁶ Fe ⁺ intensity	δ ⁵⁶ Feraw (‰)	δ ⁵⁶ Feirmm014 (‰)	2SE
Balmat-3nA@22	6.13E+07	-33.50	-0.38	0.09
Balmat-3nA@23	6.09E+07	-33.50	-0.38	0.10
Balmat-3nA@24	5.97E+07	-33.60	-0.48	0.08
Balmat-3nA@25	6.23E+07	-33.53	-0.41	0.10
Balmat-3nA@26	6.17E+07	-33.47	-0.35	0.10
Balmat-3nA@39	6.11E+07	-33.39	-0.23	0.08
Balmat-3nA@40	6.18E+07	-33.52	-0.36	0.11
Balmat-3nA@41	6.07E+07	-33.38	-0.22	0.10
Balmat-3nA@42	6.27E+07	-33.45	-0.29	0.10
Balmat-3nA@43	6.33E+07	-33.77	-0.61	0.09
Balmat-3nA@44	6.21E+07	-33.66	-0.50	0.09
Balmat-3nA@45	6.32E+07	-33.75	-0.59	0.07
Balmat-3nA@50	6.63E+07	-33.97	-0.63	0.09
Balmat-3nA@51	6.31E+07	-33.65	-0.30	0.11
Balmat-3nA@53	6.12E+07	-33.57	-0.22	0.09
Balmat-3nA@54	6.47E+07	-33.79	-0.44	0.08
Balmat-3nA@55	6.28E+07	-33.90	-0.55	0.08
Balmat-3nA@56	6.32E+07	-33.61	-0.26	0.09
Balmat-3nA@57	6.51E+07	-33.44	-0.50	0.09
Balmat-3nA@58	6.48E+07	-33.11	-0.18	0.09
Balmat-3nA@59	6.57E+07	-33.31	-0.38	0.08
Balmat-3nA@60	6.53E+07	-33.29	-0.36	0.08
Balmat-3nA@61	6.66E+07	-33.51	-0.58	0.10
Balmat-3nA@62	6.37E+07	-33.35	-0.41	0.07
Balmat-3nA@63	6.28E+07	-33.45	-0.51	0.13
Balmat-3nA@64	6.26E+07	-33.19	-0.25	0.09
Balmat-3nA@65	6.09E+07	-33.26	-0.32	0.10
Balmat-3nA@66	6.30E+07	-33.43	-0.49	0.09
Balmat-3nA@67	6.58E+07	-33.34	-0.41	0.07
Balmat-3nA@69	6.64E+07	-33.50	-0.47	0.09
Balmat-3nA@70	6.44E+07	-33.48	-0.46	0.10

Balmat-3nA@71	6.73E+07	-33.35	-0.32	0.09
Balmat-3nA@72	6.63E+07	-33.37	-0.35	0.10
Mean IMF (‰)		-33.10		
Mean Repro (‰,				
2SD)		0.39		
Repro. bracketing (‰, 2SD)		0.24		

898 TABLE S3 56 Fe⁺ intensity, raw and IMF-corrected ${\delta}^{56}$ Fe values associated with Balmat 899 pyrite standard analyses (February session in CRPG-IPNT, 57 Fe/ 54 Fe ratios were not acquired 900 during this session). The internal uncertainties are given as 2SE.