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1 **Experimental investigation of Ru isotope fractionation between metal,**
2 **silicate and sulfide melts**

3

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18

19 **Abstract**

20 To improve the understanding of large-scale planetary processes, i.e. differentiation and core
21 formation, of Earth and other planetary bodies, we performed experiments at 1 GPa in a range of

22 temperatures to investigate mass-dependent isotope fractionation of ruthenium (Ru) between metal,
23 silicate, and sulfide melts. Metal silicate fractionation is $^{102}\text{Ru}/^{99}\text{Ru}_{\text{silicate}} - ^{102}\text{Ru}/^{99}\text{Ru}_{\text{metal}} = 0.02 \pm 0.02$
24 ‰ (95% confidence interval) at 1600 °C and therefore negligible for Earth's core formation. However,
25 there is resolvable Ru isotope fractionation between liquid metal and liquid sulfide: The $^{102}\text{Ru}/^{99}\text{Ru}$
26 ratio of liquid sulfide is 0.11 ± 0.03 ‰ lighter than that of liquid metal at 1400 °C in sulfur (S)-bearing
27 experiments. The unexpected lighter Ru isotope composition of the sulfide can be best explained with
28 different Ru-S bonding environments. Our results show furthermore, that addition of tin (Sn) instead
29 of S to experimental charges affects Ru isotope fractionation significantly. The $^{102}\text{Ru}/^{99}\text{Ru}$ ratios in the
30 Sn-bearing phase are $0.18 \text{ ‰} \pm 0.01 \text{ ‰}$ heavier than metal; hence, the presence of Sn not only changes
31 the magnitude of the Ru isotope fractionation but also its direction. The observed Ru isotope
32 fractionations are too small to preserve a resolvable isotope fractionation signature during core
33 formation or the Hadean matte scenario at very high temperatures.

34

35 **1. Introduction**

36 Planetary core formation, i.e., the segregation of a metal core from a molten silicate mantle is one of
37 the most fundamental processes during planetary differentiation. The highly siderophile elements
38 (HSE – Ru, Rh, Pd, Re, Os, Ir, Pt, Au) are expected to strongly partition into Earth's metallic core during
39 its formation. However, the partitioning behavior of some HSE may depend on the conditions of core
40 formation. Many experimental studies determined metal-silicate partition coefficients for the HSE and
41 suggest that Earth's mantle should have been left more depleted in most HSEs than it is actually
42 observed (e.g. O'Neill et al., 1995; Borisov and Palme, 2000; Holzheid et al., 2000; Fortenfant et al.,
43 2003; Ertel et al, 2008; Mann et al., 2012). Moreover, the relative chondritic HSE abundances inferred
44 for Earth's mantle cannot easily be explained with metal-silicate partitioning. This discrepancy
45 between the expected and actual HSE abundances in Earth's mantle led to the 'late veneer hypothesis'
46 which invokes a late accretion of chondritic material to Earth's mantle after core formation ceased
47 (e.g., Kimura et al., 1974; Wänke 1981; Walker et al., 2015). In consequence, the elemental and isotopic

48 composition of the HSE in the Earth's mantle should reflect predominantly the composition of the
49 material that was added after core formation.

50 A promising tool to better understand core formation and late accretion is stable isotope fractionation
51 during core formation: Recent experimental and theoretical studies have shown that stable isotopes
52 of several elements, e.g. Cr, Fe, Mo, Si and Sn, are significantly fractionated between liquid metal and
53 silicate phases (e.g. Roskosz et al., 2006; Georg et al., 2007; Roskosz et al., 2009; Moynier et al., 2011;
54 Shahar et al., 2011; Hin et al., 2012, 2013, 2014; Shahar et al., 2015; Bourdon et al., 2018). Mass
55 dependent isotope fractionation of an element is controlled by differences in the bonding environment
56 of the element in different phases (Bigeleisen and Mayer, 1947; Urey, 1947); therefore, stable isotope
57 fractionation is often observed due to different oxidation states or coordination of the element in
58 different phases (Bourdon et al., 2018). As the magnitude of equilibrium isotope fractionation is
59 strongly dependent on temperature ($\sim 1/T^2$) (Bigeleisen and Mayer, 1947; Urey, 1947), stable isotope
60 data may be useful to constrain the temperature of metal-silicate equilibration and therefore improve
61 our understanding of core formation processes (review about various isotopic systems and methods
62 in Bourdon et al., 2018). The isotope fractionation factor can potentially be calculated ab-initio from
63 vibrational frequencies of the isotopes in different phases (Bigeleisen and Mayer, 1947; Urey, 1947),
64 but as these remain poorly constrained for most elements, high-pressure high-temperature
65 experiments have been employed to determine mass-dependent stable isotope fractionation between
66 metal and silicate phases (Roskosz et al., 2006; Georg et al., 2007; Roskosz et al., 2009; Shahar et al.,
67 2011; Hin et al., 2012, 2013, 2014; Shahar et al., 2015).

68 In this study, we conducted experiments to investigate mass-dependent Ru isotope fractionation
69 between liquid metal and liquid silicate. Ru plays an important role in the understanding of late
70 accretion: e.g. natural peridotites show elevated Ru/Ir relative to chondrites (Becker et al., 2006; Day
71 et al., 2017). So far, this was either explained by the addition of a fractionated iron meteorite-like
72 component during late accretion (Fischer-Gödde and Becker, 2012), or by sulfide segregation during
73 Earth's core formation, as Ru is less chalcophile than Ir or other HSE at high pressure and high

74 temperature (Laurenz et al., 2016; Rubie et al., 2016). However, the process that caused elevated Ru/Ir
75 has not yet been identified. Hopp and Kleine (2018) compared the mass-dependent Ru isotopic
76 signatures of chondrites, iron meteorites, and terrestrial peridotites and concluded that the bulk
77 silicate Earth (BSE) has a chondritic mass-dependent Ru isotope signature and fractionated iron
78 meteorite components could not have been added during late accretion. Furthermore, the observation
79 of a chondritic Ru isotopic composition of the BSE predicts that core formation processes did not
80 induce measureable Ru isotope fractionation in Earth's mantle prior to the addition of a chondritic late
81 veneer (Hopp and Kleine, 2018). However, experimental constraints on the direction and magnitude
82 of Ru isotope fractionation between metal and silicate are required to evaluate if any significant
83 amount of Earth's mantle Ru could derive from pre-late veneer processes, i.e. core formation or sulfide
84 segregation. In detail, experimental proof of significant Ru isotope fractionation during core formation
85 processes would, in combination with the chondritic Ru isotopic composition of the BSE, exclude any
86 contribution of such isotopically fractionated pre-late accretion Ru to Earth's mantle HSE budget.

87 Experiments of this study were designed to study Ru isotope fractionation between metal and silicate.
88 As we will show below, many Ru isotope analyses of silicate phases in this study are dominated by the
89 formation of sulfide nuggets caused by the experimental setup, hence, display metal-sulfide isotope
90 fractionation instead of metal-silicate fractionation. The interaction of metal and sulfide affects HSE
91 during differentiation of planetary bodies, i.e. partial melting and core crystallization, and can
92 therefore give insights into the evolution of two meteorite groups: (1) Magmatic iron meteorites are
93 remnants of asteroidal cores that underwent fractional crystallization (e.g. Scott, 1972; Scott and
94 Wasson, 1975). Hopp et al. (2018) showed that with progressive fractional crystallization the
95 crystallizing solids become isotopically heavier. This can be explained either by the extraction of
96 isotopically lighter Ru in early formed S-poor solids that led to increasingly heavier isotopic
97 composition in the remaining S-rich melt or by a kinetic effect during fractional solidification of the
98 metallic melt. (2) The HSE systematics of samples from partial differentiated asteroids, i.e. primitive
99 achondrites, are controlled by interaction of metal and sulfide phases that form during the early stages

100 of differentiation (e.g. Goodrich et al., 2013; Keil, 2014; Keil and McCoy, 2018). Hence, experimental
101 constraints on Ru isotope fractionation between metal and sulfide potentially can provide insights into
102 early differentiation processes.

103

104 **2. Methods**

105 **2.1 Starting Materials**

106 The starting material was prepared from analytical grade oxides (SiO_2 , Al_2O_3 , MgO , FeO), carbonates
107 (CaCO_3 , Na_2CO_3 , K_2CO_3), pure S and metals (Fe, Sn, Ru). To release absorbed water and unwanted
108 hydroxides, MgO was fired in a Pt-crucible to 1000 °C for at least 2 h and it was subsequently stored in
109 a drying oven at 110 °C. The starting material compositions are listed in Table 1. We always used the
110 same batch of Ru powder for the preparation of all starting materials. Silicate as well as metal powders
111 were mixed separately in agate mortars under acetone for at least 30 min to obtain fine grained and
112 homogenous starting material mixtures.

113 The silicate starting material was first prepared without Fe. This Fe-free mixture of oxides and
114 carbonates was vitrified in a Pt crucible at 1500 °C for 2h to release all CO_2 from the starting material.
115 The glass was reground under acetone and FeO was subsequently added to the silicate starting
116 powder.

117 As Ru partition coefficients can easily exceed values of 10000 (Laurenz et al., 2016), extremely low Ru
118 concentrations are expected for the silicate glass. Therefore, 10 to 11 wt.% Ru was added to the metal
119 phase starting material. Assuming a metal-silicate ratio of 1:10 and a $D_{\text{Ru}}^{\text{metal-silicate}}$ of 10 000, this results
120 in a silicate glass with 1.1 $\mu\text{g/g}$ Ru.

121 **2.2 Experimental techniques**

122 The starting material was pressed into either MgO single crystal or graphite capsules (Schunk,
123 Germany). MgO capsules were drilled out of 10 mm³ synthetic MgO single crystal cubes in the technical

124 workshops at the Institute for Mineralogy at WWU Münster. In order to maximize the capsule size and
125 the amount of silicate glass in the run products, the capsule size was set to 6 mm O.D., 4 mm I.D., with
126 an inner length of about 6 mm and a 2 mm thick bottom. The capsule was closed by a 2 mm thick lid
127 of the same capsule material.

128 Silicate and metal starting material powders were pressed into the capsule in separate layers with a
129 volume ratio of about 90 % silicate and 10 % metal. For most of the experiments, the metal layer was
130 placed on top of the silicate powder to ensure that the metal melt sinks through the entire column of
131 silicate melt during the experiment. However, to test for any effect of the metal location in the capsule
132 before the runs on isotope fractionation, several experiments were performed where the metal
133 powder was located at the bottom of the capsule, and other runs (e.g. E017) contain starting materials
134 where metal and silicate mixtures were homogeneously mixed (Table 3).

135 All experiments were conducted in an end-loaded piston cylinder apparatus at the University of
136 Münster using $\frac{1}{2}$ " piston cylinder assemblies consisting of concentric cylinders of talc, pyrex glass and
137 a thin graphite heater (Klemme et al. 1997). The inner crushable alumina parts of our usual piston-
138 cylinder assemblies (e.g. Gervasoni et al., 2017; Grützner et al., 2017) were replaced completely by the
139 single crystal MgO capsules or were reduced to 0.5 mm thickness when graphite capsules were used.
140 This modification of the piston cylinder assembly allows experiments at relatively high pressures and
141 provides enough material in a single experimental run for Ru isotope analyses. Temperatures were
142 monitored and controlled with $W_{97}Re_3$ - $W_{75}Re_{25}$ thermocouples and a Eurotherm-controller (Schneider
143 Electric).

144 The experimental run durations ranged from 2 to 6 h. The experiments were run for a minimum of 2h
145 to achieve chemical and isotopic equilibrium between liquid metal and liquid silicate. This is commonly
146 achieved in minutes to < 2h (e.g. Poitrasson et al., 2009; Shahar et al., 2011; Hin et al., 2012, 2013,
147 2014; Bourdon et al., 2018) for related systems. Sample E098 and E102 show the same isotope
148 fractionation values within the error for 2h and 6h which shows that isotopic equilibrium is reached at

149 least for 1600 °C in less than 2h. The same is the case for < 3h at 1300 °C (sample E104 and E128).
150 However, given the case of isotopic disequilibrium in a certain experiment we would expect higher
151 fractionation than at equilibrated conditions. As we will show in the Discussion, fractionation is small.
152 It would then become even smaller at equilibration without changing the proposed implications of this
153 study. The run duration was not longer than 6h to avoid any Soret effect that may cause isotope
154 fractionation along the temperature gradient of the piston cylinder capsule but would normally require
155 several tens of hours run duration (Kyser et al., 1998; Richter et al., 2008).

156 As equilibrium isotope fractionation strongly depends on temperature and decreases significantly with
157 increasing temperature, experiments were performed between 1300 and 1600 °C. The lower
158 temperature limit was set by the liquidus temperature of the metal phase and the upper limit by the
159 technical limitations of the piston cylinder, hence by the amount of needed material. In order to lower
160 the liquidus of the metal phase, Sn was added to the metal phase in the first runs (c.f. Hin et al., 2013).
161 However, Young et al. (2015) and Shahar et al. (2015) argue that addition of Sn to the metal might
162 affect the isotopic fractionation. Therefore, we replaced Sn by S in other runs, or we used a mixture of
163 both S and Sn (Table 1).

164 The experimental run products were either mounted in epoxy or acrylic resin for electron microscopy,
165 electron microprobe, and Laser ablation inductively coupled mass spectrometry measurements, or the
166 capsules were carefully crushed in an agate mortar for Ru isotope analysis. Due to the material loss
167 during mounting, cutting, and polishing the mounted samples had not enough material for successful
168 further Ru isotope analyses. Therefore, most samples were either mounted for element concentration
169 analyses with the techniques mentioned above, or they were prepared for isotope analysis.

170 For isotope analysis the silicate glass was separated from the capsule rim and from the metal both
171 using hand picking and magnetic separation. If possible, a representative but randomly picked piece of
172 silicate glass was chosen to be mounted in epoxy resin for additional elemental analysis using EPMA
173 and LA-ICPMS (Table 2).

174 Experiment E135 was first mounted in epoxy for element concentration analysis. Afterwards, a small
175 core (< 4 mm) of the silicate glass was drilled out. The capsule walls with attached remaining silicate
176 glass were cut in several cross sections to study Ru concentrations inside the capsule and metal, as
177 well as sulfide nugget distribution along the capsule walls.

178 **2.3 Major element analysis**

179 The experimental run products were first characterized by optical microscopy, followed by scanning
180 electron microscope imaging and energy-dispersive X-ray spectroscopy (EDS) analysis (JEOL 6510LA).
181 To quantify major and minor element concentrations of the experimental products, a JEOL 8530F
182 electron microprobe (EPMA) was used. Measurement conditions for wavelength-dispersive
183 spectroscopy (WDS) were typically 15 keV acceleration voltage, 5 nA beam current and usually 10 μm
184 beam diameter to receive representative analyses. Standards were pure metals for metal composition
185 and synthetic silicates and oxides for the basaltic glass. S was standardized on pyrite for metals and on
186 celestine for silicates, respectively. Due to a potential peak shift, peak searches were done for S before
187 the analysis. Ru concentrations in silicates were always below the detection limit.

188 Investigations of the observed nanonuggets were conducted using semi-quantitative EDS element
189 mapping in and quantitative WDS analysis. As for the small droplet size of ≤ 400 nm the accelerating
190 voltage was set to 7kV. At such conditions Monte-Carlo simulations (Drouin et al., 2007) show a lateral
191 resolution of about 400 nm enabling quantitative analyses of the few larger 400 nm sized melt
192 droplets. However, as revealed by apparent Si concentrations of about 5 wt.% a small portion of the
193 surrounding silicate glass has also been excited.

194 WDS analyses were performed in two steps. Firstly, all elements except Ru and Sn were analyzed with
195 a beam current of 15 nA and counting times of 15 s on the peak and 5 s on the background. Secondly,
196 the same spots were measured for Ru and Sn at 100 nA and counting times of 80 s on the peak and 40
197 s on the background. Finally, both analyses were merged using the Offline-matrix correction provided
198 by the JEOL instrument software. Silicon, S ($K\alpha$ line) and Fe, Sn, Ru ($L\alpha$ line) were standardized on the
199 large metal phase within the same sample (10 μm spot size) to avoid variations in carbon coating

200 thickness. The metal phase elemental composition had been quantified before under “normal”
201 conditions as described above.

202 **2.4 Trace elements**

203 Trace elements were analyzed using a Laser ablation inductively coupled mass spectrometry system
204 (Analyte G2 Excimer laser with a wavelength of 193 nm coupled to a Thermo Scientific Element 2 HR-
205 ICPMS) at the Institute for Mineralogy at WWU Münster. The laser was operated with a repetition rate
206 of 10 Hz and an energy fluence of 3 J/cm² per pulse. Typical spot sizes were 65 µm. The ablated material
207 was transported to the ICP torch using He and Ar as carrier gases with flow rates of 0.13 L/min.
208 Measurements on the reference material (NIST 612 glass standard) were done at the beginning and
209 end of every session, as well as after about every 30 spots. All analyses were normalized to FeO
210 concentrations, which had been determined by EPMA analyses. For internal reference and to track any
211 mass bias drift, we used BIR-1G, BHVO-2G, and metal phases from experimental run E017 and E031.
212 The analytical results were processed using the software GLITTER (Griffin et al., 2008).

213 **2.5 Ruthenium isotope analysis**

214 The Ru stable isotopic composition of the experimental run products were analyzed following the Ru
215 double spike method described by Hopp et al. (2016). The small sample weights, the large amount of
216 Ru in the metal phases, as well as the contrasting low and unknown Ru concentration of the silicate
217 glass required modification of the procedures that are described in the following paragraph.

218 **2.5.1. Sample preparation**

219 After separation of metal and crushing of silicate glasses, the samples were weighed and transferred
220 into Savillex PFA beakers. Metal phases (~0.02 to 0.05 g) were digested using 7.5 ml *reverse aqua regia*
221 inside sealed Carius tubes at 220 °C for 2 days (Shirey and Walker, 1995). These metal phases always
222 contain the bulk of the Ru in the experiment (~10 wt.%) and hence represent the bulk/initial Ru isotopic
223 compositions of the system. To verify this, an aliquot of the starting material (Alfa Aesar Ru > 99.9 %
224 metal basis powder; 325 mesh, 13994, LOT: R17A033) was digested along with the metals and analyzed

225 for its Ru isotopic composition. After digestion, a 1 ml aliquot of the solutions were diluted to 10 ml of
226 6 M HCl.

227 The silicate samples (~0.06 to ~0.13 g) were digested in PFA vials at 100 °C for 24 hours using a mixture
228 of double teflon-distilled HCl and HF acids (4 ml concentrated HCl + 2 ml HF). After digestion, aliquots
229 of the metal and silicate sample solutions were converted to 0.28M HNO₃ and the Ru concentrations
230 in the solutions were determined using a Thermo Scientific *X-Series 2* quadrupole ICPMS at the Institut
231 für Planetologie, Münster. Based on the calculated Ru concentrations in the individual samples an
232 appropriate amount of a ⁹⁸Ru-¹⁰¹Ru double spike was added. This double spike is used to correct for all
233 analytical fractionation effects; thus, precise determination of the natural mass-dependent isotope
234 fractionation is possible (Rudge et al., 2009). The spiked solutions were equilibrated in closed beakers
235 for 24 hours using *reverse aqua regia* at 120 °C on a hotplate. This procedure results in complete spike-
236 sample equilibration, as demonstrated for iron meteorites and chondrites (Hopp et al., 2018; Hopp
237 and Kleine, 2018). Additionally, no differences between meteorite samples that were spiked prior or
238 after digestion were observed (Hopp and Kleine, 2018). Furthermore, digestion and sample-spike
239 equilibration of several powder aliquots of a homogeneous ~100 g powder of the Allende meteorite
240 using the method described above resulted in reproducible and accurate data (Hopp and Kleine, 2018).

241 2.5.2. Chemical separation of Ruthenium

242 The chemical separation of Ru followed a two-stage ion exchange chromatography procedure based
243 on the method outlined in Hopp and Kleine (2018). After sample-spike equilibration, the solutions were
244 dried down at 100 °C on a hotplate, converted to chloride form and re-dissolved in 5 ml of 0.2 M HCl.
245 In the first step, the sample solutions were loaded onto cation exchange columns filled with 10 ml pre-
246 cleaned BioRad AG50 W-X8 (100-200 mesh) resin. On these columns, the bulk of the HSE were eluted
247 in a total volume of 14 ml 0.2 M HCl, while the major elements (i.e., Fe and Ni) remain adsorbed on
248 the resin. Then the Ru fractions were dried down on a hotplate and re-dissolved three times using 5
249 ml of 1 M HF. To remove remaining interfering elements (Zr, Mo, Pd) the Ru fractions were dissolved
250 in 7 ml 1 M HF and were loaded onto anion exchange columns filled with 2 ml of pre-cleaned BioRad

251 AG1-WX8 (100-200 mesh) resin. Ruthenium is eluted in 14 ml 1 M HF, whereas Zr, Mo and Pd remain
252 adsorbed onto the resin. The final Ru fractions were dried and re-dissolved in 0.5 ml 0.28 M HNO₃. This
253 procedure results generally in pure Ru fractions with Mo/Ru < 0.0005 and Pd/Ru < 0.001, which are
254 sufficiently low to allow for reliable correction of interferences on ⁹⁸Ru, ¹⁰⁰Ru, ¹⁰²Ru and ¹⁰⁴Ru, i.e. the
255 isotopes used in the double spike inversion (Hopp et al., 2016). The overall yield of the chemical
256 separation of Ru varies between ~60 and ~90 %. The Ru blank is 49 ± 33 pg Ru (1 s.d.; n = 5) and hence
257 insignificant, given that more than 70 ng Ru was analyzed for each sample.

258 2.5.3. Mass spectrometry and data reduction

259 The Ru isotope measurements were conducted using a Thermo Scientific Neptune *Plus* MC-ICPMS at
260 the Institut für Planetologie at University of Münster. Prior to the measurements, the samples were
261 dissolved in 0.28M HNO₃ and were introduced into the mass spectrometer using a CETAC *Aridus* II
262 desolvating system combined with a 70 µl/min ESI PFA nebulizer. The formation of oxides was
263 monitored as CeO/Ce and reduced to < 1 % by the addition of N₂ to the sample gas. Sample and
264 standard solutions were measured at ~100 ppb using conventional Ni H cones and ion beams were
265 simultaneously collected in static mode for all seven stable Ru isotopes (⁹⁶Ru, ⁹⁸Ru, ⁹⁹Ru, ¹⁰⁰Ru, ¹⁰¹Ru,
266 ¹⁰²Ru, ¹⁰⁴Ru) together with ⁹⁷Mo and ¹⁰⁵Pd as interference monitors. Ruthenium masses were measured
267 using Faraday cups connected to 10¹¹ Ω feedback resistors and the ion beams at ⁹⁷Mo and ¹⁰⁵Pd were
268 collected using 10¹² Ω feedback resistors. Spiked sample measurements comprised 50 × 8.2 s
269 integrations of the ion beams and consumed ~70 ng Ru. The baselines were measured on peak with
270 40 × 4.2 s integrations on a solution blank prior each measurement.

271 The Ru isotopic data are reported relative to the composition of an Alfa Aesar standard solution (RuCl₃;
272 Lot # 61300952) as follows:

$$273 \delta^{102/99}\text{Ru} = \left(\frac{{}^{102/99}\text{Ru}_{\text{sample}}}{{}^{102/99}\text{Ru}_{\text{standard}}} - 1 \right) \times 10^3 \quad (1)$$

274 Data reduction and calculation of $\delta^{102/99}\text{Ru}$ values was performed off-line as described in Hopp et al.
275 (2016), using either the 'double spike toolbox' (Rudge et al., 2009) or the geometrical approach

276 described in Siebert et al. (2001). Both data reduction schemes were used in this study and yielded
277 identical results. The molar proportions of spike determined with the reduction schemes were used to
278 calculate Ru concentrations in the samples. Furthermore, the data reduction provides the natural
279 fractionation factor α , from which the mass-dependent Ru isotopic composition of a sample is
280 calculated as follows:

$$281 \quad \delta^{102/99}\text{Ru}_{\text{meas.}} = -1000 \times (\alpha_{\text{sample}} - \alpha_{\text{standard}}) \times \ln(m_{102}/m_{99}) \quad (2)$$

282 where m_{102} and m_{99} are the atomic weights of ^{102}Ru and ^{99}Ru and α_{standard} is the mean composition
283 measured for the spiked Alfa Aesar standard solution in each analytical session. The external
284 reproducibility of the Ru stable isotope measurements was estimated to be ± 0.05 ‰ (2 s.d.; $n = 20$)
285 for $\delta^{102/99}\text{Ru}$, based on repeated measurements of spiked Ru standard solutions and Ru doped
286 geochemical reference materials (BHVO-2, BCR-2, UB-N) (Hopp et al., 2016) and by seven replicate
287 digestions of an Allende (CV3) chondrite powder (Hopp and Kleine, 2018).

288 Fractionation factors between metal and silicate or sulfide, respectively ($\Delta^{102/99}\text{Ru}_{\text{metal-silicate}} =$
289 $\delta^{102/99}\text{Ru}_{\text{metal}} - \delta^{102/99}\text{Ru}_{\text{silicate}}$), are calculated based on the average isotopic compositions of eleven
290 metals ($\delta^{102/99}\text{Ru}_{\text{bulk}}$) that equals the bulk isotopic composition of the starting material because > 99.99
291 % of Ru is incorporated in the metal phases.

292

293 **3. Results**

294 Representative major element compositions from SEM-EDS and EPMA measurements are listed in
295 Table 2. Samples that were only analyzed by SEM-EDS are marked in Table 2. The Ru isotopic
296 compositions of experimental products together with Ru concentrations in silicates are presented in
297 Table 3. For both EPMA and LA-ICPMS measurements a minimum of 10 spots were analyzed and used
298 for the mean values in Table 2 and Table 3.

299 **3.1 Textures**

300 *3.1.1 Silicate phases*

301 During the experiments, the melt composition changes, as the melt reacts with the MgO capsule and
302 becomes more picritic. Consequently, a thin layer of olivine crystallizes during the run at the contact
303 zone between MgO capsule and melt (Figure 1). The quenched silicate melt is typically glass with no
304 quench crystals or other phases. The experiments at 1600 °C result in elongated, needle-shaped olivine
305 crystals (Figure 2), caused by rapid dissolution of MgO in the basaltic melt (Hin et al., 2013). MgO
306 concentrations in silicate glasses at 1600 °C reach up to 36.4(2) wt.%, whereas in experiments at $T <$
307 1600 °C the MgO concentration ranges from 7.5(2) to 15.2(6) wt.% (Table 2).

308 *3.1.2 Metal blobs, metal droplets and sulfidic nanonuggets*

309 In successful experiments, a large metal blob is located at the bottom of the capsule after the run. This
310 is independent from the location of the metal powder in the capsule before the run, which was either
311 on top or at the bottom of the capsule (Figure 1a). In experiments with a homogenized starting material
312 (runs E017, E029, E031) several medium-sized blobs ($> 10 \mu\text{m}$) were found in the capsule attached to
313 the capsule walls (Figure 1a).

314 Tiny metal droplets ($< 10 \mu\text{m}$) are observed at the capsule walls in nearly all experiments (Figure 1d).
315 However, the number of droplets increases, if the metal had been placed at the top of the capsule
316 before the run and sank down to the bottom during the experiment. Droplets attached to the wall can
317 be also found in run products where the metal powder had been placed directly at the bottom.

318 Large blobs, medium blobs and tiny metal droplets are all assigned to the metal phase during the
319 experimental run and show no variation in chemical composition. Most medium size blobs are
320 attached to the olivines that formed at the capsule rim (Figure 1). Their number can be minimized by
321 placing the starting metal material at the bottom of the capsule.

322 Beside the aforementioned metal phases, nano-sized ($< 1 \mu\text{m}$) nuggets appear in many silicate glasses
323 (Figure 6) and these nuggets have a composition that differs strongly from the metal starting material.

324 The analyzed nuggets are all sulfidic and because of their small size could not be separated from the
325 silicate phase. As it will be shown in subsection 4.1.2, these Ru-bearing sulfidic nuggets are in
326 equilibrium with the melts during the experiments and they dominate the budget and isotopic
327 composition of Ru in the silicate portion of the experiments. In the following text, the term ‘nugget’
328 hence refers to the sulfidic nanonuggets described above. Any small metal phase of similar size that
329 has a composition identical to the major metal phase in the experiment or any Fe-exsolution phase
330 that formed during quenching, will be labeled as ‘droplet’.

331 In graphite capsule experiments, we did not observe metal droplets attached to the walls or sulfidic
332 nanonuggets. This correlates well with the lack of olivine in experiments with graphite capsules. These
333 results indicate that metal droplets do not stick to the capsule wall itself, but that they tend to attach
334 to surfaces of crystallizing olivines close to the wall of MgO capsules (Figure 1).

335

336 3.2 Oxygen fugacity

337 The oxygen fugacity was mainly controlled by the reducing starting material components. To get an
338 idea of the oxygen fugacity in different experiments it was calculated using the equation

$$339 \Delta IW = 2 \times \log \left(\frac{X_{FeO} \cdot \gamma_{FeO}}{X_{Fe} \cdot \gamma_{Fe}} \right) \quad (3)$$

340 where X_{FeO} and X_{Fe} are the molar fractions of iron in the silicate and metal melts. γ_{FeO} and γ_{Fe}
341 describe their respective activity coefficients. The activity coefficient for Fe in the metal alloy was
342 calculated using the modified Wagner epsilon formalism (Ma, 2001). Interaction parameters for Sn and
343 S were taken from the steel making literature (JSPS, the 19th Committee on Steelmaking, 1988). The
344 addition of up to 4 wt.% S and/or 23 wt.% Sn to the Fe-dominated alloy shows almost ideal behavior.
345 For Ru an ideal approach to Raoult’s law was assumed due to the lack of activity data. The calculated
346 values for γ_{Fe} range from 0.995 to 1.067. For γ_{FeO} the value was set to 1.7 after Holzheid et al. (1997),
347 who showed that the activity coefficient of FeO in a basaltic melt with MgO between 4-20 wt.% and

348 FeO up to 12 wt.% is constant between 1300 °C and 1600 °C. If we exclude the experiments at 1600 °C
349 with MgO concentrations of above 30 wt.%, all other silica glasses from this study are within the
350 aforementioned range (Table 2). In MgO capsules oxygen fugacity vary between $\Delta IW = -1.86$ and -2.05 .
351 This is in good agreement with the approximation made by Wade and Wood (2005) for their proposed
352 core formation conditions. In graphite capsules oxygen fugacity is expected to be in a similar range
353 around IW but fO_2 could not be calculated in these experiments as the metal phase was lost during
354 sample preparation. In these experiments the metal phase did not form a large metal blob, but instead
355 a metal layer was formed that was also attached to the capsule wall. However, the existence of the
356 metal phase indicates an oxygen fugacity of at least $\Delta IW < 0$ in these runs.

357 **3.3 Ruthenium concentrations in experimental run products**

358 Ruthenium concentrations were analyzed with EPMA or SEM-EDS for metal phases and with LA-ICPMS
359 as well as MC-ICPMS for the silicate phases. Results for Ru in metal and silicate analyses are listed in
360 Table 3. In the metal phases Ru concentrations range from 10.8(2) to 14.0(8) wt.%. The coexisting
361 sulfide phase from E041 has a Ru element concentration of 0.7(2) wt.%. Ruthenium concentrations
362 measured with LA-ICPMS in silicate glasses range from 0.02 $\mu\text{g/g}$ to 0.6 $\mu\text{g/g}$. The only exception is
363 sample E029 with a Ru concentration of 8 $\mu\text{g/g}$ in the glass. Where the entire capsule was embedded
364 in epoxy for LA-ICPMS analyses (E017; E135), laser spots were chosen to form a cross section along the
365 capsule from the bottom to the top. No Ru concentration gradient was found in the silicate glass of
366 these samples, and we conclude that Ru is homogeneously distributed in the silicate glass (Figure 3).

367 MC-ICPMS analyses show generally higher concentrations of Ru in the silicate fractions than LA-ICPMS
368 analyses. This is caused by the (sulfidic) equilibrium nanonuggets, which could not be separated from
369 the silicate phase during preparation (except for experiment E135 – see below). Glasses from graphite
370 capsule experiments have the lowest Ru concentrations measured with MC-ICPMS (7 to 8.2 $\mu\text{g/g}$).
371 These concentrations are still higher than the LA-ICPMS data, however, in the range of concentrations
372 determined in the glass of E029. Silicate fractions from MgO capsule experiments have Ru

373 concentrations ranging from 9.7 $\mu\text{g/g}$ up to 199 $\mu\text{g/g}$ when measured with MC-ICPMS – up to three
374 orders of magnitude higher than the results from LA-ICPMS.

375 Most runs were prepared for MC-ICPMS analysis by crushing of the sample and separating the metal
376 from the silicate phase – either by picking or by magnetic separation with a hand magnet. For
377 experiment E135 the entire capsule was mounted in epoxy for EPMA and LA-ICPMS analyses. Laser
378 spots were set on the glass, the capsule walls, on the core of olivine and at olivine rims (Figure 2). The
379 LA-ICPMS data show that the Ru concentrations in the glass and in olivine are extremely low (< 0.1
380 $\mu\text{g/g}$) while at the olivine surface ($> 0.1 \mu\text{g/g}$) and at the capsule rim ($\gg 1 \mu\text{g/g}$) the Ru concentrations
381 are elevated (Figure 4a). In the MgO capsule Ru concentrations are below the detection limit. After the
382 LA-ICPMS analyses, the silicate glass was drilled out of the mounted capsule with a miniature core drill
383 to avoid any contamination with capsule material or olivine crystals from the wall. Additionally, the
384 olivine-rich glass at the contact to the metal was separated and the MgO capsule was cut into four
385 different sections (Figure 4b) and analyzed in the same way as described for the glasses. Ru
386 concentrations of all seven samples including the metal phase and a zone of ol-rich glass can be found
387 in Figure 4b. Ru concentrations of the silicate glass are low ($0.4 \mu\text{g/g}$) and in good agreement with LA-
388 ICPMS analyses. Besides the metal phase, only the ol-bearing glass ($5.4 \mu\text{g/g}$) and the lowermost
389 capsule wall close to the metal phase ($158 \mu\text{g/g}$) show elevated Ru concentrations. The analyzed
390 capsule wall sections show Ru concentrations from $0.04 \mu\text{g/g}$ to $0.28 \mu\text{g/g}$ in the three upper sections
391 (B, C, D in Figure 4b). The metal phase contains 14.0(8) wt.% Ru.

392 **3.4 Ruthenium isotope compositions**

393 The Ru isotopic composition of the eleven analyzed metal phases ranges from $\delta^{102/99}\text{Ru}$ of 0.02 ‰ to
394 0.07 ‰ (average of $\delta^{102/99}\text{Ru}_{\text{metal}} = 0.04 \pm 0.02 \text{ ‰}$ (95 % conf.), without any clear trends related to the
395 starting material, temperature or capsule material (Table 3). This average is in good agreement with
396 Ru isotopic composition of the starting material of $\delta^{102/99}\text{Ru} = 0.03 \pm 0.02 \text{ ‰}$ (95 % conf.; $n = 4$). Thus,
397 as predicted from the highly siderophile character of Ru, the metals in our runs are very close to the
398 bulk Ru isotopic composition of the experiments.

399 The Ru isotopic signatures in the silicate phases show some fractionation trends depending on the
400 chemical system (Figure 5): The silicate glasses in S-bearing experiments (E088, E089, E097, E098 and
401 E102) have isotopically lighter Ru compositions with $\delta^{102/99}\text{Ru}$ values ranging from $-0.01 \pm 0.01 \text{ ‰}$ to -
402 $0.07 \pm 0.03 \text{ ‰}$. In contrast, the silicate portions of the two Sn-bearing experiments E110 and E126 have
403 isotopically heavier Ru compositions, with $\delta^{102/99}\text{Ru}$ values of $0.22 \pm 0.01 \text{ ‰}$ and $0.21 \pm 0.01 \text{ ‰}$,
404 respectively. The three experiments that contain both Sn and S show only small or no Ru isotope
405 fractionation, with Ru isotopic compositions of the silicate portions varying between $\delta^{102/99}\text{Ru} = -0.01$
406 $\pm 0.02 \text{ ‰}$ and $0.06 \pm 0.01 \text{ ‰}$. Moreover, the silicate glasses of the experiment without S or Sn (E134)
407 has a Ru isotopic composition of $\delta^{102/99}\text{Ru} = 0.02 \pm 0.02 \text{ ‰}$ which is very close to the bulk Ru isotopic
408 composition of the metal phases. Similarly, the S-bearing experiments conducted in graphite capsules
409 (E115, E141) show no Ru isotopic fractionation, with silicate glasses having a $\delta^{102/99}\text{Ru}$ of 0.03 and 0.05
410 ‰ , respectively.

411

412 **4. Discussion**

413 **4.1 Ruthenium concentrations in silicate glasses**

414 Ruthenium concentrations determined by double spike MC-ICPMS are generally 2-4 orders of
415 magnitude larger than those obtained by LA-ICPMS. Whereas the Ru concentrations in the silicate
416 glasses measured with LA-ICPMS are close to the detection limit, MC-ICPMS analyses return
417 concentrations of up to 200 $\mu\text{g/g}$ Ru. Both analytical methods give good and reproducible results for
418 all analyzed reference materials (Methods). Hence, analytical errors, that cause up to three orders of
419 magnitude difference, can be excluded. The high concentrations in the silicate fractions of the
420 experiments determined by MC-ICPMS were first expected to be most likely a result of either external
421 contamination during sample preparation or of insufficient physical separation of silicate glass and Ru-
422 bearing nuggets or droplets. The latter has been seen previously for Ir, Pt and Re (O'Neill, et al., 1995;
423 Ertel et al., 2008; Medard et al., 2016). Both scenarios were not the case as we show in the following.

424 4.1.1. Testing for metal contamination of the silicate glass

425 The possibility of external contamination was evaluated with a metal-free experiment (E125) that was
426 performed and processed in the same way as all former experiments. The Ru concentration measured
427 with LA-ICPMS and quadrupole ICPMS was below the detection limit for both methods and hence
428 excludes any external contamination of the samples during preparation for MC-ICPMS analysis. To
429 exclude contamination by incomplete separation of metal and silicate phase after the experiment (e.g.
430 small metal droplets in the silicate or at the capsule wall), experiment E135 was conducted and
431 analyzed as described above. Ruthenium concentrations in the silicate measured by LA-ICPMS are low
432 (0.02 $\mu\text{g/g}$) and LA-ICPMS profiles show no gradients across the silicate glass (Figure 3). Analyses of
433 olivine phenocrysts that settled at the bottom of the capsule close to the main metal phase show also
434 no enrichment in Ru (Figure 2). However, the olivine surface (i.e., the olivine-glass interface) as well as
435 analyses at the capsule rim show elevated Ru concentrations of up to 1 $\mu\text{g/g}$, which is about two orders
436 of magnitude higher than the Ru concentrations in the glass (Figure 4a). Optically visible, but i.e. μm -
437 sized, metal droplets were clearly too small to be analyzed with LA-ICPMS. We analyzed the droplets
438 together with surrounding material (Laser spot size: 20-60 μm) and the results show Ru concentrations
439 from 50 to 500 $\mu\text{g/g}$. Further analyses were done on E135 glass in the center of the capsule and on
440 glass of other samples to ascertain that the glass itself is nugget- and droplet-free. Using LA-ICPMS,
441 elevated Ru concentration can be only found at the capsule walls attached to olivines (Figure 1d).

442 These observations can be best explained by Ru-bearing sulfide nuggets or metal droplets that are
443 attached to the olivine phenocrysts in the ol-rich glass and to the newly formed olivines at the capsule
444 wall. As the metal starting material was placed at the bottom of the capsule in run E135, Ru-bearing
445 nuggets stayed attached only in lower sections of the capsule wall, during melting and during
446 formation of the large metal phase. No Ru-rich nuggets were transported upwards and stuck to capsule
447 wall sections in the top part of the capsule. Olivine itself should normally store small amounts of Ru.
448 Capobianco et al. (1991) and Brenan et al. (2003) find $D_{\text{Ru}}^{\text{olivine-silicate}}$ between 2.2 and 0.2, depending on

449 f_{O_2} and T . However, $D_{Ru}^{silicate-sulfide}$ is very small (c.f. section 4.2) and we can expect similar values for
450 $D_{Ru}^{olivine-sulfide}$, which agrees well with the lack of measurable amounts of Ru in olivine.

451 If we assume that the Ru-rich nuggets at the capsule walls are droplets from the primary metal, they
452 should have the same isotopic signature as the main metal phase. In contrast, the measured Ru isotope
453 compositions of the nugget-containing silicate portions are systematically fractionated relative to the
454 metal. Hence, contamination cannot be caused by the observed metal droplets at the capsule wall.
455 Existing metal droplets are part of the primary metal and can neither explain the good reproducibility
456 of the fractionated $\delta^{102/99}Ru$ values (Table 2) nor the lack of correlation between Ru concentration and
457 $\delta^{102/99}Ru$ values (Figure 3). However, this is the case for sulfide nuggets.

458 *4.1.2. Formation of sulfidic nanonuggets*

459 The nanonuggets in experiment E176 are attached to the basalt glass-olivine interface and have a
460 diameter of about ≤ 400 nm (Figure 6). The Ru concentration in the nuggets was too low to be mapped.
461 Quantitative WDS analyses give Ru concentrations of about 435 ± 120 $\mu g/g$ which are far above Ru
462 concentrations in the S-doped silicate, but they are not fully reliable on nuggets of this size as all
463 analyses resulted in mixed analyses (Supplementary Table 1). This is caused by contamination of the
464 surrounding basaltic glass and olivine (too large beam excitation volume). Of about 15 analyzed
465 nuggets only two gave totals ≥ 98 wt.% and ≤ 102 wt.%. Nevertheless, all analyses show strongly
466 increased S concentrations of about 25-35 wt.% (Supplementary Table 1). Therefore, we assume that
467 the nanonuggets have a sulfidic composition and represent droplets of a third immiscible liquid that
468 formed during the experiment. The generation of larger amounts of sulfide melt in coexistence with
469 metal and silicate phase is shown in experiment E041 (Figure 2). The sulfide layer around the metal
470 phase has 27(1) wt.% S, 64.0(8) wt.% Fe and 0.7(2) wt.% Ru. Assuming similar Ru concentrations in
471 sulfide nuggets, 400 to 7000 $\mu g/g$ are far above Ru concentration in basalt glass. The formation of these
472 sulfidic nuggets can explain the discrepancy between Ru concentrations determined by LA-ICPMS and
473 MC-ICPMS analyses, as well as the excellent reproducibility of $\delta^{102/99}Ru$ values among S-bearing

474 experiments in MgO capsules. Moreover, the lack of a correlation between Ru concentrations and
475 isotopic compositions shows that MC-ICPMS results are dominated by the sulfide nanonuggets and
476 that significant contributions of initial metal droplets can be excluded.

477 To add 13 $\mu\text{g/g}$ Ru to the silicate melt bound in sulfide nuggets (with about 500 to 7000 $\mu\text{g/g}$ Ru), only
478 a small contribution of sulfide nuggets to the separated silicate fractions is required. For example,
479 based on the masses and Ru concentrations of the silicate sample E098 it can be estimated that a
480 sulfide nugget contribution of 3.2 wt.% to 0.2 wt.%, respectively, is sufficient to explain the observed
481 high Ru concentrations in the MC-ICPMS analysis.

482 Sulfide nuggets produced in metal-silicate partition experiments have been already reported by Shahar
483 et al. (2015) who describe “submicrometer sized iron sulfides” within their silicate glass that are in
484 equilibrium with the glass. We assume that the sulfidic nuggets form during melting of the metal phase
485 at the beginning of the experiment. Their attachment to olivines and to the capsule wall argues against
486 late exsolution of Ru-rich nuggets from the melt during quenching. Quench nuggets would be
487 dispersed over a larger range within the glass and would be detectable due to elevated Ru
488 concentrations by LA-ICPMS (or a better agreement of LA-ICPMS and MC-ICPMS results). Experiment
489 E110 (Figure 1b) shows such exsolution textures after quenching, however the exsolution droplet
490 composition is pure Fe-metal. As the silica melt from which these quench nuggets precipitated can
491 only store very small amounts of Ru, LA-ICPMS analyses of E110 glass and quench metal droplets
492 cannot contain any elevated Ru concentrations (0.3(1) $\mu\text{g/g}$).

493 To explain the change in Ru isotope fractionation between S- and Sn-bearing experiments, equivalent
494 Sn-rich nuggets could exist in the Sn-rich, S-free experiments (E110; E126). Fe-rich and Sn-rich phases
495 may coexist at temperatures > 1400 °C (Franke et al., 2005). Unfortunately, Sn-rich nuggets could not
496 be detected in our experiment with EPMA in “low keV mode”. Instead of Sn nuggets, one could also
497 speculate about an alternative scenario: as FeSn alloys have relatively low melting points, larger
498 amounts of Sn could soften the bonding environment in the molten alloy. If Ru would have a bonding

499 affinity to Sn in the alloy, light Ru isotope bonding would be promoted leaving the silicate isotopically
500 heavier. However, this would require the Ru isotope signature to be derived by the silicate and not by
501 Sn-nuggets. In this case, Sn-nuggets are not required and a low Ru concentration would be expected
502 for the bulk silicate analysis with MC-ICPMS, which is not the case for Sn-bearing experiments (e.g.
503 198.7 $\mu\text{g/g}$ in E126).

504 This section showed that the analyzed $\Delta^{102/99}\text{Ru}$ of most experiments is dominated by either a S- or Sn-
505 rich phase in the silicate glass. The following sections will further discuss the unambiguous nature of
506 the Ru isotopic composition analyzed in the silicate glass, the impact of varying experimental
507 conditions and the mechanisms that might drive these effects.

508

509 **4.2 Ruthenium metal-silicate partition coefficients**

510 There is scant data on the Ru partitioning between metal and silicate melts or sulfide and silicate melts.
511 Ruthenium element partition coefficients can be calculated in this study from Ru concentrations in the
512 metal and LA-ICPMS analyses of the glasses. For S-bearing experiments $\log D_{\text{Ru}}^{\text{metal-silicate}}$ range from
513 2.0(1) to 6.8(3). A $\log D_{\text{Ru}}^{\text{sulfide-silicate}}$ value of 4.6(2) was calculated for experiment E041. Varying amounts
514 of S in the metal starting materials (0.001 wt.%; 0.01 wt.%; 0.1 wt.%) show that $\log D_{\text{Ru}}^{\text{metal-silicate}}$
515 decreases with increasing amount of S in the system (Figure 7). This is in good agreement with
516 experimental studies from Laurenz et al. (2016). They show that Ru and other HSEs become less
517 siderophile with increasing S concentration in the silicate melt. The calculated $\log D_{\text{Ru}}^{\text{metal-silicate}}$ and \log
518 $D_{\text{Ru}}^{\text{sulfide-silicate}}$ from Laurenz et al. (2016) are slightly lower when compared to values from this study
519 (Figure 7). The different partition coefficients may be explained by different pressures or
520 temperatures, as Laurenz et al. (2016) reported that $\log D_{\text{Ru}}^{\text{metal-silicate}}$ decrease with increasing pressure
521 and our runs were performed at 1 GPa, whilst experiments from Laurenz et al. (2016) were conducted
522 at 11 GPa (Figure 7).

523 4.3 Ruthenium isotope fractionation

524 Mass-dependent Ru isotope fractionation between quenched metal melt and the nugget-bearing
525 silicate glasses is small but detectable (Figure 5). Sulfur-bearing experiments in MgO capsules show an
526 average Ru isotope fractionation factor $\Delta^{102/99}\text{Ru}_{\text{metal-sulfide}}$ of $0.11 \pm 0.03 \text{ ‰}$ (95 % conf.) where
527 $\Delta^{102/99}\text{Ru}_{\text{metal-sulfide}} = \delta^{102/99}\text{Ru}_{\text{metal}} - \delta^{102/99}\text{Ru}_{\text{sulfide}}$. The two Sn-bearing experiments in MgO capsules show
528 $\Delta^{102/99}\text{Ru}_{\text{metal-Sn-phase}}$ values of $-0.18 \text{ ‰} \pm 0.01 \text{ ‰}$. These differences in the direction of the isotope
529 fractionation can be best explained with the sulfidic or and Sn-rich phase, respectively, that dominate
530 the isotopic signature. The very small isotope fractionation in the S- and Sn-free experiments in a MgO
531 capsule (E134; Table 3) shows that in the S- or Sn-bearing experiments the measured Ru isotope
532 fractionation is controlled by formation of S- or Sn-bearing phases, respectively. The S+Sn-bearing
533 experiments show an average Ru isotope fractionation factor $\Delta^{102/99}\text{Ru}_{\text{metal-sulfide}}$ of $-0.02 \pm 0.02 \text{ ‰}$
534 (Figure 5). These runs were originally conducted to lower the melting temperature and allow
535 experiments at 1300 °C. Given these low fractionation factors, it could be assumed that sulfide
536 nanonuggets and a Sn-phase exists in the silicate glass. The two effects of S- and Sn bearing phases
537 would counterbalance each other in experiment E104, E109 and E128 and we would expect a
538 correlation between S- and Sn-content in the silicate phase. However, both elements were not
539 measured with MC-ICPMS and the samples could not be analyzed by EPMA. The Ru concentration in
540 the silicate glass is also not higher (22-96 $\mu\text{g/g}$) than in experiments with only S or Sn in the starting
541 material.

542 In graphite capsules olivine cannot form at the capsule walls and hence the formation of S-rich nuggets
543 is prohibited. Thus, the Ru concentration in the silicate fractions of the graphite experiments are low
544 (7-8 ppm, measured with MC-ICPMS) and reproducible. Nevertheless, these concentrations are higher
545 than results from LA-ICPMS measurements and the formation of S-rich nuggets, like in MgO-capsule
546 experiments, can be excluded due to the lack of any Ru isotope fractionation in graphite experiments.
547 The slightly higher Ru concentration in silicates from graphite capsule experiments can be best
548 explained by contamination with primary metal droplets.

549 For experiments at 1600 °C Hin et al. (2013) found that isotope fractionation starts to be dominated
550 by kinetic fractionation in MgO capsules as the element of interest (Mo in their study) fractionates
551 from the metal phase into a silicate whose composition changes over time. Therefore, Hin and
552 coworkers treated these experiments separately. At 1600 °C the MgO capsule dissolves much faster
553 and Hin et al. (2013) see an increase of Mo element concentration, as well as a change of isotope
554 fractionation over time. Ruthenium experiments at 1600 °C in MgO capsules from this study also show
555 significant increase of MgO in the melt, extensive olivine growth (Figure 2), and metal exsolution within
556 the silicate melt. However, LA-ICPMS data from E098 and E102 show no elevated Ru concentrations.
557 They show comparable isotope fractionation to experiments conducted at temperatures < 1600 °C.
558 Thus, the change in silicate melt composition does not affect the Ru concentration in the silicate melt.

559 **4.4. Ruthenium bonding environment and its influence on isotope fractionation**

560 Our experiments are not designed to study Ru element coordination in the melt or its valence state in
561 the different phases. Discrete data are lacking but element bonding is crucial for isotope fractionation
562 and will be discussed shortly.

563 Ru has a higher valence state in the silicate melt than in the metal melt. While the Ru oxidation state
564 in the metal is 0, Ru can be +II, +III, +IV, +VI or +VIII when bonded to non-metals. Laurenz et al. (2013)
565 propose Ru^{4+} to be the dominant cationic species for their experiments in picritic melts. In their
566 experiments, Ru bonds either with O^{2-} or S^{2-} , albeit the bonding with S^{2-} is about 900 times stronger
567 due to the chalcophile character of Ru (Laurenz et al., 2013). For RuS_2 in sulfide nanonuggets, the
568 valence state of Ru would vary between metal and sulfide from 0 to +IV as it is supposed to be for
569 metal and silicate. Ru isotope fractionation should be dominated by the varying bonding length of Ru-
570 Fe, Ru-O and Ru-S. Schauble (2004) shows that higher valence states and lower coordination states are
571 generally associated with stiffer bonds. These stiff bonds are favored by the heavier isotope of the
572 element of interest (Bigeleisen, 1965). However, this does not seem to be the case for the sulfide
573 nanonuggets, and the isotope fractionation observed in our S-bearing experiments in MgO capsules.
574 All analyzed sulfide nuggets are isotopically lighter than the metal and therefore predict a stronger

575 affinity of lighter Ru isotopes for the sulfide nuggets relative to the metal phase. This is contrary to
576 expectations from theory (Bigeleisen, 1965; Schauble, 2004) and cannot be simply explained by a lower
577 valence state than +IV. The isotopic fractionation of other elements such as Fe in S-bearing systems is
578 also not well constrained. Several studies (Williams et al., 2006; Schuessler et al., 2007; Polyakov and
579 Soultanov, 2011) show that Fe sulfides are depleted in heavy Fe isotopes relative to their silicates. In
580 contrast, Shahar et al. (2015) find good correlations between increasing sulfur content in the metal
581 and higher $\Delta^{57}\text{Fe}_{\text{metal-silicate}}$ values, which they conclude is caused by different bonding structures of
582 sulfides and S-rich metal alloys. Their results show that the heavier Fe isotopes have a stronger affinity
583 to bond with S in sulfides but not with S in metal. Hence, one could also speculate that other factors
584 like coordination of Ru or Fe within the melts have a stronger effect on isotope fractionation than their
585 valence state.

586 For the Sn-bearing experiments of this study, it can be concluded that the negative $\Delta^{102/99}\text{Ru}$ values are
587 derived from coordination differences and a Ru-Sn bonding that varies from the sulfide phase, as well
588 as from the Fe dominated metal phase. However, Sn can theoretically have valence states between –
589 IV and +IV but not much is known about Ru bonding in such phases. The effect of Sn on metal isotope
590 fractionation in experiments has been addressed in a few studies. Shahar et al. (2015) and Young et al.
591 (2015) propose that large amounts of Sn in the metal melt changes the bonding environment and
592 should therefore influence isotope fractionation of elements such as Fe or Mo. Nevertheless, Hin et al.
593 (2013) conducted experiments on Mo isotope fractionation between Sn-free and Sn-bearing metal and
594 silicate melts, and could not find any significant differences. In our experiments, there is a significant
595 effect of Sn (in the metal phase) on Ru isotope fractionation (c.f. also Hin et al. 2012; 2014). But to
596 better understand Ru isotope fractionation between liquid metal, sulfide and silicate, further work is
597 required on Ru valence state, bonding environment, and coordination in these phases.

598 **4.5 Implications for Ru isotope fractionation in natural Fe-FeS systems**

599 There is no clear temperature dependency in Figure 5 for metal-sulfide fractionation experiments
600 between 1400 °C and 1600 °C as their variation is too small. To better describe the change of Ru

601 isotope fractionation with varying temperature a best-fit approach can be applied: Bigeleisen and
602 Mayer (1947) and Urey (1947) show that equilibrium mass dependent isotope fractionation is
603 proportional to A/T^2 , where A is a constant that can be either determined from experiments or
604 theoretically (e.g. Bourdon et al., 2018). This results in a decreasing magnitude of isotope fractionation
605 with increasing temperature. To extrapolate the effect of temperature on Ru isotope fractionation,
606 one can use a Least Square Approach for A/T^2 as it is shown in Figure 8. For this fit a value of $0.24 (\pm$
607 $0.03) \times 10^6$ was calculated for A on the base of the S-bearing experiments at 1400 °C, 1500 °C and 1600
608 °C (E088, E089, E097, E098, E102; Table 3). Thus, the temperature dependence of the Ru isotope
609 fractionation between metal and sulfide can be described as:

$$610 \quad \Delta^{102/99}Ru_{\text{metal-sulfide}} = \frac{0.24(\pm 0.03) \cdot 10^6}{T^2} \quad (6)$$

611 The fractionation of $\Delta^{102/99}Ru_{\text{metal-sulfide}}$ is not resolvable ($<0.04 \text{ ‰}$) at temperatures above 2500 °C
612 (Figure 8b) and $\Delta^{102/99}Ru_{\text{metal-silicate}}$ is already negligible at $> 1600 \text{ °C}$. Given these results from Figure 8
613 and the even smaller silicate-metal fractionation ($0.02 \pm 0.02 \text{ ‰}$ at 1600 °C), significant Ru isotope
614 fractionation during Earth's core formation ($T > 3500 \text{ °C}$) is not expected. These findings are in good
615 agreement with the results from Hopp and Kleine (2018) who did not see any distinguishable Ru
616 isotope fractionation between chondritic meteorites and terrestrial peridotites.

617 It has been recently shown that the partitioning behavior of some HSEs may be different under Earth's
618 core formation conditions and that they will not be quantitatively removed from the mantle by metal-
619 silicate partitioning but would be overabundant and display non-chondritic relative abundances
620 (Righter et al., 2008; Rubie et al., 2016; Righter et al., 2018). Therefore, the relative chondritic
621 abundances of HSEs in Earth's mantle require an additional removal of HSEs by a sulfide matte during
622 the late stages of core formation before a chondritic late veneer is added (O'Neil, 1991; Rubie et al.,
623 2016; Righter et al., 2018). This Hadean matte model proposes that the exsolution of FeS from the
624 magma ocean during cooling affects HSE/Ir ratios in the Earth's mantle, as they show different behavior
625 (Laurenz et al., 2016; Rubie et al., 2016). However, high temperatures above 2000 °C (rather even 3000

626 - 4000 °C) are also expected for the Hadean matte (Savage et al., 2015), and according to our findings
627 for metal-silicate and metal-sulfide fractionation, significant sulfide-silicate Ru isotope fractionation is
628 also not expected at these temperatures.

629 The observed metal-sulfide isotope fractionation of Ru stable isotopes could be important to
630 understand planetary processes occurring at lower temperatures that are closer to the temperatures
631 from our experiments: Hopp et al. (2018) determined mass-dependent Ru isotope signatures of
632 various magmatic iron meteorites. These meteorites stem from the metallic cores of differentiated
633 protoplanetary bodies. The magmatic iron displays heavy and variable $\delta^{102/99}\text{Ru}$ compared to
634 undifferentiated material such as ordinary chondrites. The $\delta^{102/99}\text{Ru}$ values vary between -0.07 ± 0.05
635 and 0.85 ± 0.05 and show a correlation of increasingly heavier Ru isotopic composition with decreasing
636 Ru contents. Hopp et al. (2018) interpret these findings with progressive fractional crystallization of
637 the metallic liquid, which leads to enrichment in FeS and heavier Ru isotope signatures in the remaining
638 melt (Hopp et al., 2018). These results suggest that fractionation of Ru was controlled by the increasing
639 amount of sulfur in the melt which might cause Ru isotope fractionation between Fe and FeS
640 dominated phases. This fractionation model requires that the remaining S-rich melt is isotopically
641 heavier than the solidifying metal, which is contrary to results from this study. However, the phases
642 involved during the crystallization of magmatic iron meteorites are S-poor solid metal and S-rich
643 metallic metal that are distinct to the S-rich metal and sulfide nuggets that control the Ru isotopic
644 fractionation in our experiments. Furthermore, the incongruent crystallization itself might play a role
645 in the Ru isotope fractionation of magmatic iron meteorites but is not represented in the experiments.
646 Hence, the experimental results on the Ru isotope fractionation do not represent the mechanisms that
647 control Ru isotope fractionation during fractional crystallization of asteroidal cores.

648 Primitive achondrites are meteorites that derive from partially differentiated parent bodies (e.g.,
649 Weisberg et al., 2006). Thus, while magmatic iron meteorites are expected to have already experienced
650 core formation and core mantle segregation on their parent body, primitive achondrites represent
651 earlier stages of differentiation. These meteorites experienced only partial silicate melting and

652 incomplete metal segregation (e.g. Keil, 2014; Keil and McCoy, 2017). Nevertheless, they provide
653 evidence for variable extraction of S-rich and S-poor Fe-Ni-S melts (Day et al., 2012; Goodrich et al.,
654 2013; Dhaliwal et al., 2017; Day et al., 2019). Hopp and Kleine (2021) reported that primitive
655 achondrites from different groups, i.e. brachinites, acapulcoite-lodranites, and ureilites, have heavier
656 $\delta^{102/99}\text{Ru}$ values relative to chondrites. The extraction of S-rich Fe-Ni-S melts cannot easily explain the
657 observed Ru isotopic fractionation with reasonable degrees of melt extraction. However, the
658 extraction of S-poor metallic melts at higher temperatures can produce the heavier $\delta^{102/99}\text{Ru}$ values of
659 primitive achondrites with realistic degrees of melt extraction (Hopp and Kleine, 2021). Thus, the
660 results from the S-rich experiments reported here are likely not directly applicable to the main driver
661 of isotopic fractionation in primitive achondrites, i.e. extraction of S-poor metallic melts. However,
662 they might represent initial extraction of S-rich Fe-Ni-S during partial differentiation of planetesimals
663 that contributed to the bulk isotopic composition. Interestingly, the observed Ru isotope fractionation
664 in primitive achondrites agrees with the magnitude and direction of the isotope fractionation observed
665 in this study.

666

667 **5. Conclusions**

668 Our experiments were designed to study mass-dependent isotope fractionation of stable Ru isotopes
669 between metal melt and silicate melt. This metal-silicate fractionation turned out to be too small and
670 negligible for core formation temperatures. However, in several experiments significant Ru isotope
671 fractionation was caused by metal-sulfide fractionation with a lighter Ru isotope composition in the
672 sulfide phase. The addition of small amounts of S (used as a flux) to the experimental system led to the
673 formation of Ru-bearing sulfide nuggets which dominated the measurements. This effect could only
674 be seen in MgO capsules but not in graphite capsules. The direction of this fractionation contradicts what
675 would have been expected by theoretical approaches from literature: a heavier isotopic composition
676 for Ru in a non-metallic valence state. Moreover, the addition of Sn instead of S to the starting material
677 results very likely in a Sn-rich phase that has a heavier Ru isotope composition than the Fe-metal. In

678 consequence, the exchange of small amounts of S with Sn (or vice-versa) in the starting material can
679 flip over the direction of Ru isotope fractionation in the experimental system. There are scant data
680 about the coordination and the real valence state of Ru in sulfides or other compounds. To better
681 understand the effect of the chemical composition on Ru (or HSE in general) isotope fractionation, we
682 need to better understand their bonding in the studied phases.

683 The magnitude of Ru isotope fractionation is small in our experiments. Since equilibrium isotope
684 fractionation decreases with increasing temperature, we do not expect any resolvable isotope
685 fractionation during core formation temperatures (> 2000 °C) or a late sulfide segregation from a
686 magma ocean.

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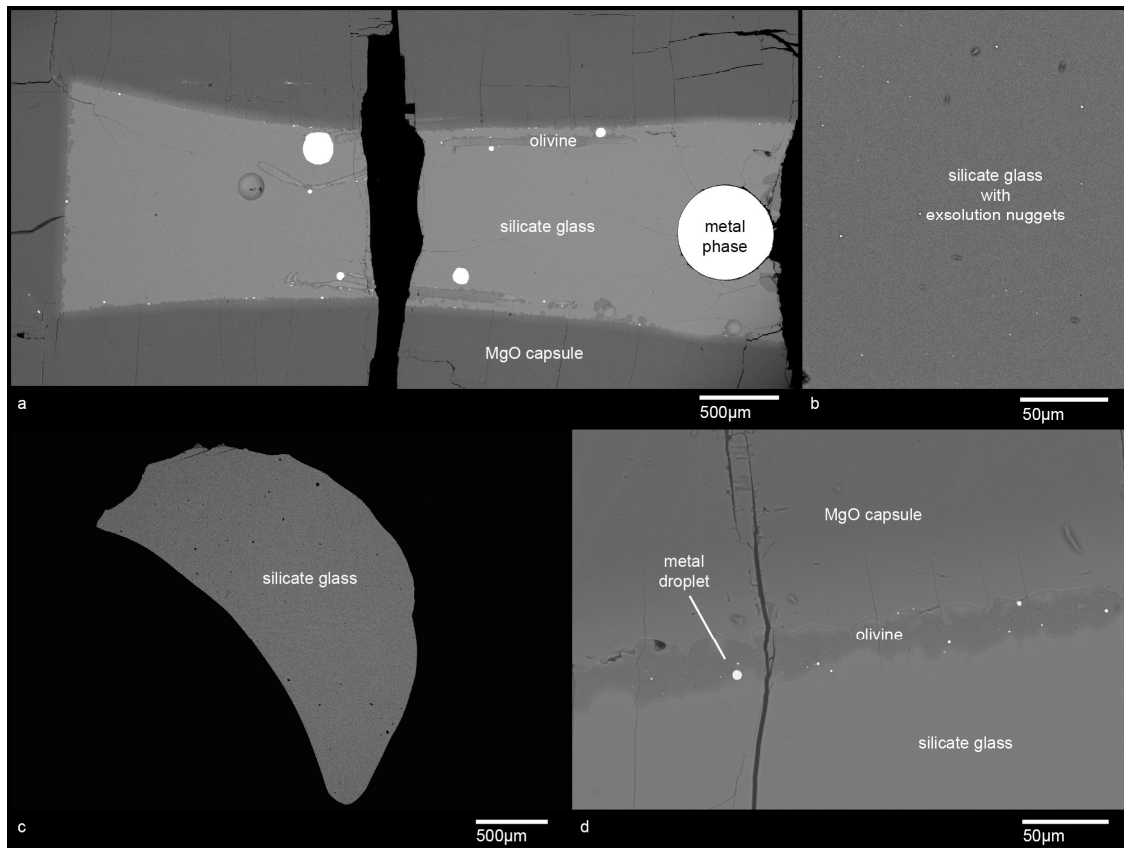
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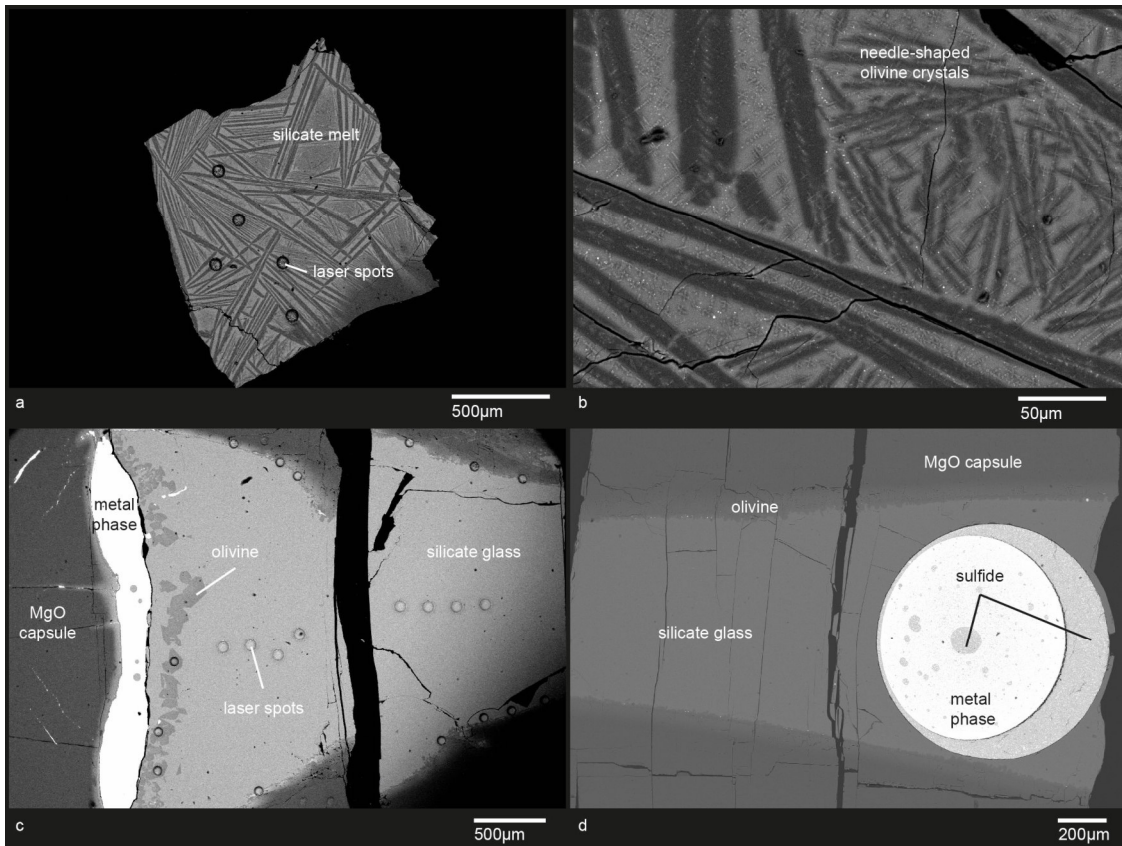
870 **Figures**



871

872 *Figure 1: Back-scattered electron (BSE) image of experiments a) E017, b), c) E110 and d) E041. a) A*
873 *homogeneous mixture of metal and silicates in the starting materials results in numerous medium*
874 *size metal blobs in the run product. b) and c) nano size nuggets (bright spots) in the silicate glass are*
875 *found in several experiments but show these nuggets results in no elevation in Ru concentrations, and*
876 *hence these nuggets are interpreted to be formed by Fe-exsolution during the quench of the run. d)*
877 *metal droplets that form during the run and that are attached to the olivine layer at the capsule wall.*

878



879

880 *Figure 2: Back-scattered electron (BSE) image of experiments a) E134, b) E102, c) E135 and d) E041.*

881 *a), b) Needle-shaped olivine textures in experiments at 1600 °C are caused by fast dissolution of MgO*

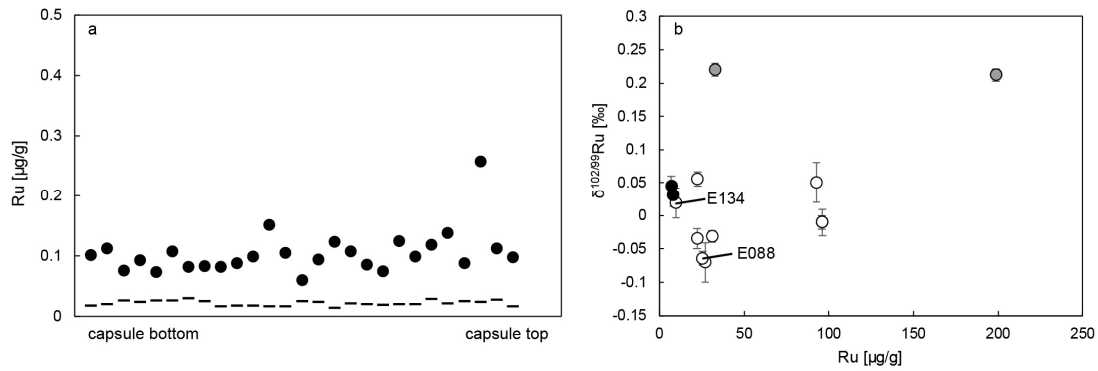
882 *in the basaltic melt. c) nano size nuggets (bright spots) are found in the silicate glass of several*

883 *experiments but show no elevation in Ru concentration and hence these nuggets are interpreted to be*

884 *formed by Fe-exsolution during the quench of the run. d) In runs with high amount of S in the starting*

885 *material, a sulfide layer forms around the metal phase.*

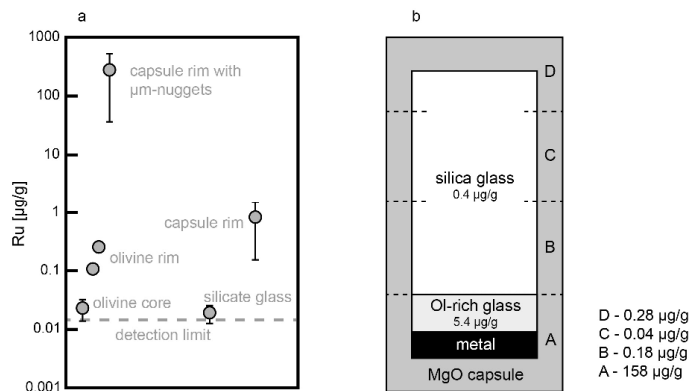
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887

888 *Figure 3: a) Ruthenium concentrations (µg/g) in the silicate measured with LA-ICPMS. Concentrations*
 889 *were analyzed along the cross sections of run E017 (circles; detection limit = bars) from the bottom of*
 890 *the capsule, i.e. close to the metal phase, to the top of the capsule (Figure 1a). Ruthenium*
 891 *concentrations in the silicate show no gradient between top and bottom of the capsule. b) Ruthenium*
 892 *concentrations (µg/g) in the silicate measured with MC-ICPMS vs. δ^{102/99}Ru in the silicate samples*
 893 *show no correlation, which is a strong argument against contamination by a metal phase in the*
 894 *analyzed silicate glass. White = S-bearing runs; grey = Sn-bearing runs; black = runs in graphite*
 895 *capsule.*

896

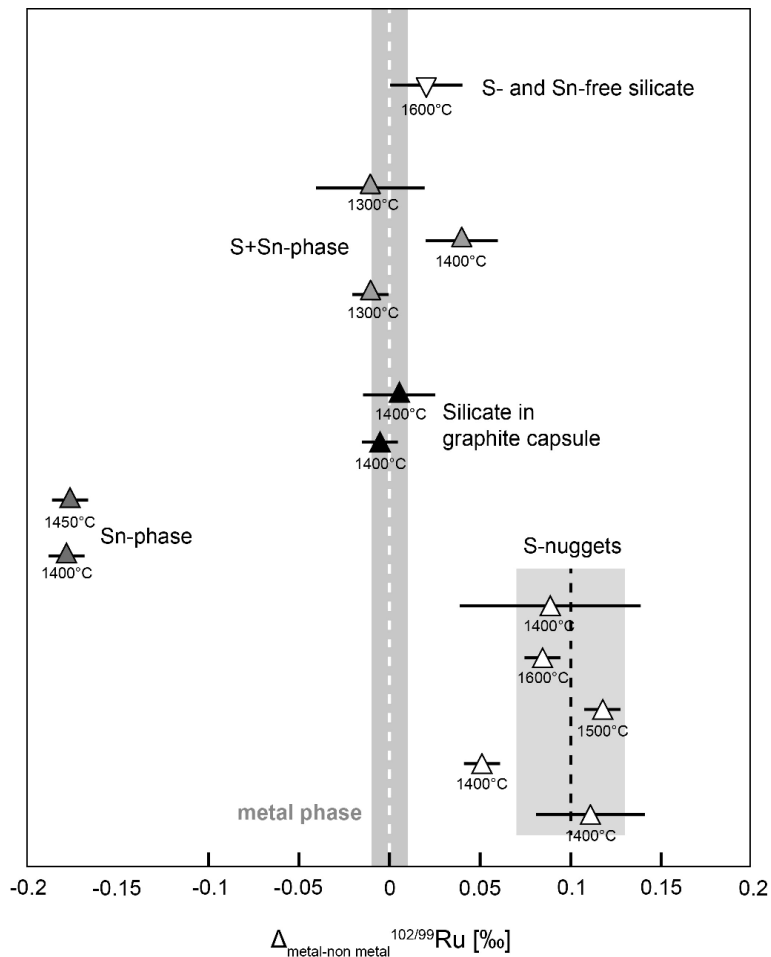


897

898 *Figure 4: Ruthenium concentrations measured in run E135. SEM image of E135 can be is shown in*
 899 *Figure 2c. a) While Ru concentrations measured by LA-ICPMS in the silicate glass and inside the*
 900 *olivine crystals are low, Ru concentrations are elevated at the rim of the olivine phenocrysts and at*
 901 *the olivine layer at the edge of the capsule. Concentrations in the MgO capsule were below the*

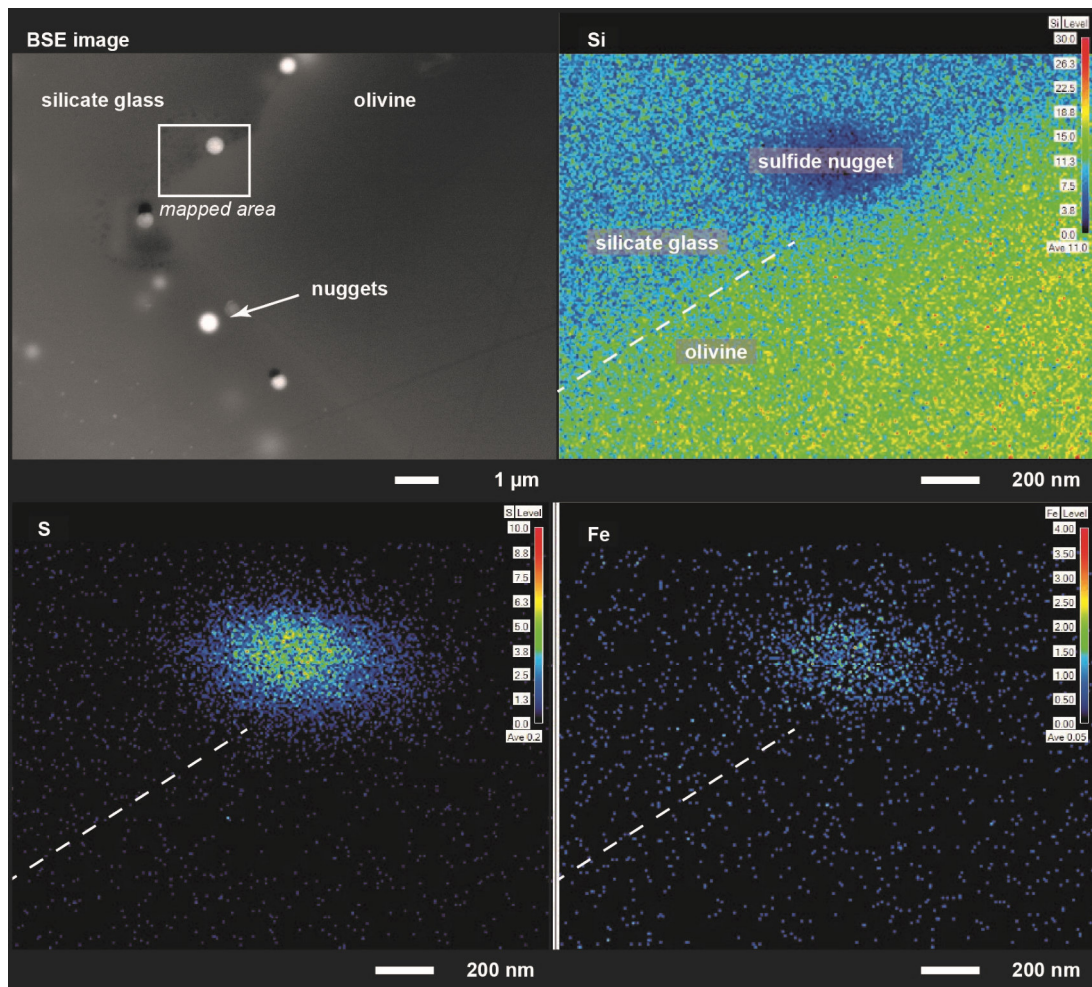
902 detection limit. This supports the assumption the Ru is concentrated in Ru-rich nuggets, which are
 903 attached to the silicate glass-olivine interface. b) Sketch of experiment E135. The sample was
 904 separated into 7 pieces, which were then separately analyzed with quadrupole ICPMS: metal, silicate
 905 glass, ol-rich silicate glass, 4 (A-D) sections of the MgO capsule: The silicate glass was drilled out of
 906 the capsule leaving a rest of glass that was attached to the capsule (section A-D). The “ol-rich” glass
 907 (with olivine phenocrysts, SEM image in Figure 2c) and the glass attached to the capsule fragments A-
 908 D show elevated Ru concentrations relative to the drilled-out silicate glass. This supports the
 909 assumption that the Ru is concentrated in Ru-rich nuggets, which are attached to the silicate glass-
 910 olivine interface.

911



913 *Figure 5: Ruthenium stable isotope fractionation measured in silicate glasses. The $\delta^{102/99}\text{Ru}$ analyses*
 914 *from metal are normalized to zero; $\Delta^{102/99}\text{Ru}_{\text{metal-non metal}} = \delta^{102/99}\text{Ru}_{\text{metal}} - \delta^{102/99}\text{Ru}_{\text{non metal}}$ with non-*
 915 *metal being either silicate glass, sulfide nanonuggets, or a Sn-phase. The fractionation is dependent*
 916 *on the starting material composition, as Ru is either concentrated in S-nuggets or a Sn-bearing phase.*
 917 *Runs in S-bearing systems show opposing isotope fractionation direction with Sn-bearing runs*
 918 *resulting in an isotopically lighter $\delta^{102/99}\text{Ru}$ signature for S-bearing runs. Sn-bearing experiments*
 919 *indicate a heavier signature. In experiments with graphite capsules neither S nor Sn nuggets formed*
 920 *and hence no isotope fractionation is observed.*

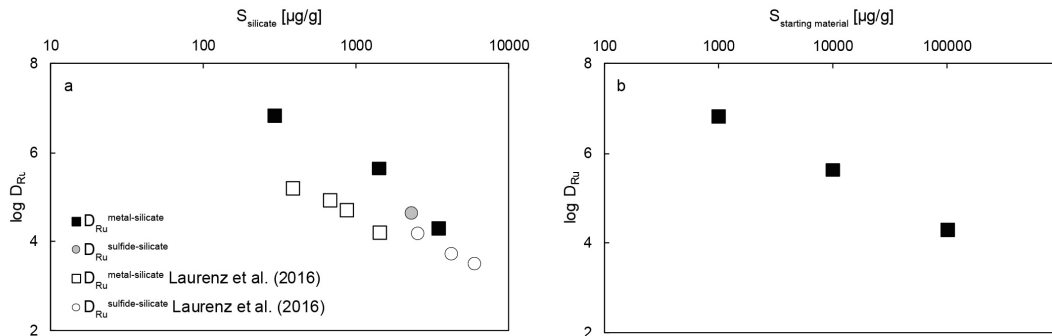
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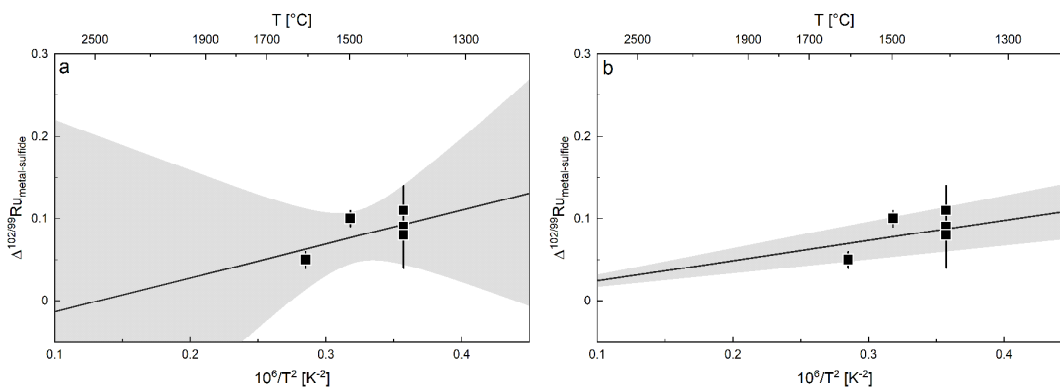
923 Figure 6: Sulfidic nanonuggets were characterized with EDS spot analysis and element mapping using
 924 EPMA. The color scale on the element maps show relative abundances of the elements. The analyzed
 925 nuggets are attached to the silicate glass-olivine interface and about 100-500 nm diameter. The white
 926 dashed line indicates the silicate glass-olivine boundary.

927



928

929 Figure 7: D_{Ru} vs. S concentrations a) in the silicate and b) in the metal portion of the starting material.
 930 Due to the presentation in logarithmic scale errors are covered within the size of the symbols. Data
 931 from Laurenz et al. (2016) are from experiments at 11 GPa and 2373 - 2473 °C. D_{Ru} increases with
 932 increasing P and T , but increasing S in the silicate makes Ru less siderophile.



933

934 Figure 8: Approach for the temperature dependence of equilibrium Ru isotope fractionation between
 935 liquid metal and liquid sulfide (E088, E089, E097, E098, E102; Table 3). Error bars are the 95 %
 936 confidence intervals of the respective analyses. a) The fit is fixed on the average value of the

937 *datapoints at 1400 °C and supports the general assumption of a temperature dependency. The*
938 *temperature dependency of the fractionation factor would be $A = 0.41 (\pm 0.33) \times 10^6$ and is*
939 *represented by the solid line. b) This weighted best fit was determined by a best fit estimation using*
940 *the average value for the 1400 °C experiments and the 1600 °C experiment and is fixed to “zero”. The*
941 *temperature dependency of the fractionation factor $A = 0.24 (\pm 0.03) \times 10^6$ is represented by the solid*
942 *line. The gray areas reflect the uncertainty (95 % confidence interval) of the fractionation factor (\pm*
943 *0.03×10^6).*

944

945

946 **Tables**

947 *Table 1 Chemical composition (wt.%) of silicate and metal starting mixtures.*

948

949 *Table 2 Major element composition of silicate glasses and metal phases in wt.% measured with*
950 *the electron probe. Uncertainties (1 s.d.) are given in parentheses showing the first significant digit.*
951 *b.d. = below detection limit. Totals indicated with * were determined by energy-dispersive X-ray*
952 *analysis (EDS) only.*

953

954 *Table 3 Details of experimental run conditions, Ru contents and Ru isotopic compositions of run*
955 *products. For some silicate samples the Ru concentrations were not only determined using bulk*
956 *methods (MC-ICPMS), but also in-situ methods using LA-ICPMS. Partition coefficients ($\log D_{Ru}^{metal-silicate}$*
957 *were calculated only for samples that were analyzed by LA-ICPMS. Capsule C = graphite, t = run duration*
958 *in hours.*

959 ^a Number of isotopic analysis.

960 ^b Ruthenium concentrations of silicate glasses determined by LA-ICPMS. Uncertainties in parentheses
961 are 1 σ of 5 to 15 spot analysis per sample.

962 ^c Ruthenium concentrations determined from the double spike isotope ratio analyses. Uncertainties
963 are <1% including measurement uncertainties and weighing.

964 ^d Measured Ru isotope mass fractionation relative to an Alfa Aesar standard solution. Given
965 uncertainties are 95% c.i. for samples with $N \geq 4$.

966 ^e Isotopic difference of metal and silicate

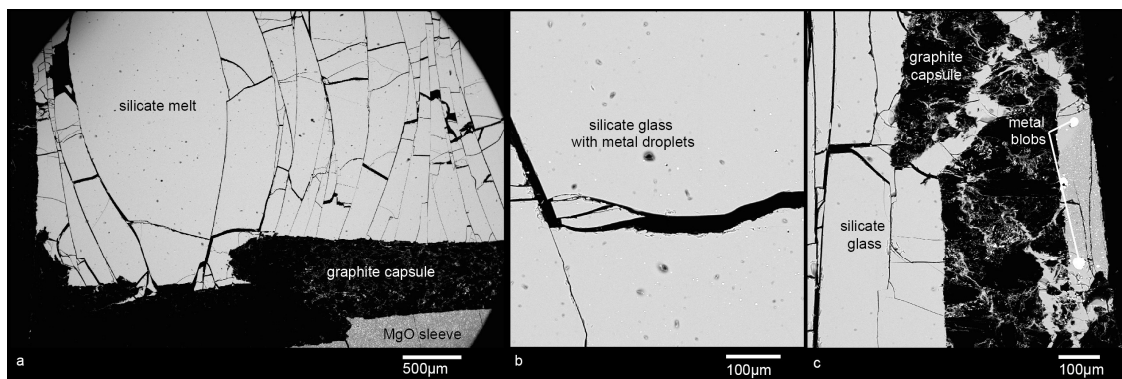
967 ^f Partition coefficients $D_{Ru}^{\text{metal-silicate}}$ calculated using LA-ICPMS concentration data of metals (Table 2)
968 and silicates for selected experiments.

969 ^g Isotopic difference of metal and silicate calculated using the average $\delta^{102/99}Ru_{\text{Bulk}}$ of all analyzed metal
970 phases representing the bulk isotopic composition as proven by the absence of any isotopic difference
971 between starting material and metal phases.

972

973

974 **Supplementary Figures**



975

976 *Supplementary Figure 1: Back-scattered electron (BSE) image of experiments E168. This experiment*
977 *was performed in a graphite capsule but could not be considered for further analyses. The metal*
978 *phase was lost during preparation. a) The contact between graphite capsule and silicate glass shows*
979 *no reaction phase like olivine in MgO capsules. b) In E168 silicate glass contains a high amount of*

980 *metal droplets that exsolved during quenching. Note that for successful runs in graphite capsule*
981 *(E115 and E141) Ru concentration in the silicate is extremely low (Table 3). c) At the bottom of the*
982 *capsule the melt partially infiltrates the the graphite capsule.*

983

Table 1: Chemical composition (wt.%) of silicate and metal starting mixtures.

	SiO ₂	Al ₂ O ₃	MgO	FeO	CaO	Na ₂ O	K ₂ O
<i>Silicate</i>							
M011	47	15	7	10	11	6.5	3.5
	Fe	Ru	Sn	S			
<i>Metals</i>							
M006	67	11	22	-			
M047	67	10	18	5			
M048	85	10	-	5			
M116	55	10	-	35			
M117	75	10	-	15			
M133	90	10	-	-			

Table 2: Major element composition of silicate glasses and metal phases in wt.% measured with electron probe micro-analysis E

Run	Na ₂ O	MgO	SiO ₂	Al ₂ O ₃	K ₂ O	CaO	S	SnO ₂
<i>Silicates</i>								
E102	3.7(2)	36.4(2)	33.7(6)	11.6(4)	2.5(1)	6.8(2)	b.d.	b.d.
E017	5.9(1)	12.6(6)	45.7(2)	15.3(3)	3.5(1)	10.0(2)	b.d.	b.d.
E110	5.8(1)	9.9(3)	47.4(2)	15.3(2)	3.2(1)	9.5(1)	b.d.	b.d.
E135	6.4(3)	12.3(3)	43.1(3)	14.7(3)	3.1(1)	10.6(2)	b.d.	b.d.
E029	5.1(2)	14.3(8)	45(1)	15.2(3)	3.0(1)	10.4(2)	0.6(1)	0.41(4)
E031	5.2(1)	15.2(7)	44.0(4)	15.1(1)	3.12(6)	10.0(3)	b.d.	b.d.
E104	5.8(3)	7.5(2)	47.6(4)	15.2(1)	3.4(1)	9.4(2)	b.d.	b.d.
E109	4.9(2)	15.7(6)	47.6(3)	13.6(7)	2.9(1)	7.3(1)	b.d.	b.d.
E134	4.0(1)	33.8(4)	34.4(8)	12.1(6)	2.5(1)	7.0(3)	b.d.	b.d.
E175	6.0(2)	7.3(3)	47.0(4)	15.3(4)	3.2(1)	9.9(2)	0.4(2)	b.d.
E041	5.6(1)	13.9(2)	43.9(3)	14.8(2)	3.3(1)	9.6(1)	0.23(3)	0.18(5)
	Fe	Si	S	Sn	Ru	Total	f(O ₂) ΔIW	
<i>Metals</i>								
E017	66.7(7)	0.06(1)	b.d.	22.9(7)	11.3(2)	101.0(3)	-1.86	
E135	81.4(8)	b.d.	4(1)	b.d.	14.0(8)	99.4(8)*	-1.86	
E031	67.4(9)	0.05(1)	0.9(2)	21(1)	10.8(2)	100.6(4)	-2.05	
E175	84.2(1)	b.d.	1.4(1)	b.d.	13.9(3)	99.5(3)*	-1.47	
E041 metal	56.2(8)	0.05(5)	5.8(3)	24(1)	13.6(2)	100.5(2)		
E041 sulfide	64.0(8)	0.01(1)	27(1)	6(1)	0.7(2)	98.2(2)		

PMA. Uncertainties (1 s.d.) are given in parentheses showing the first significant digit. b.d. = below detection limit. Totals indicated with * were

FeO	Total
5.3(2)	100.0(6)*
6.4(1)	99.5(4)
9.0(2)	100.1(3)*
9.9(2)	100.1(3)*
3.0(1)	97(1)
5.5(2)	98.3(4)
11.2(3)	100.1(4)*
8.0(3)	100.0(7)*
6.3(2)	100.1(8)*
10.9(9)	100.0(9)*
6.6(1)	98.3(4)

± determined by energy-dispersive X-ray analysis (EDX) only.

Table 3: Details of experimental run conditions, Ru contents and Ru isotopic compositions of run products. For some silicate samples the Ru c

Sample	start mixtures	T [°C]	t (h)	Capsule	Dopant	N ^a	Ru _{LA-ICPMS} [ppm] ^b	Ru _{MC-ICPMS} [ppm] ^c
<i>Starting material</i>								
Alfa Aesar Ru metal						4		
<i>Metals</i>								
E088 M	M048	1400	3	MgO	S	7		
E089 M	M048	1400	3	MgO	S	7		
E097 M	M048	1500	3	MgO	S	8		
E102 M	M048	1600	3	MgO	S	6		
E110 M	M006	1400	3	MgO	Sn	5		
E126 M	M006	1450	3	MgO	Sn	5		
E128 M	M048	1300	3	MgO	S+Sn	5		
E104 M	M047	1300	3	MgO	S+Sn	5		
E109 M	M047	1400	3	MgO	S+Sn	5		
E134 M	M133	1600	3	MgO	-	5		
E115 M	M048	1400	3	C	S	5		
<i>Average (bulk)</i>						(11)		
<i>Silicates</i>								
E088 S	M048	1400	3	MgO	S	5		26.93
E089 S	M048	1400	3	MgO	S	7		22.23
E098 S	M048	1400	6	MgO	S	5		13.70
<i>Average</i>		1400	3	MgO	S			
E097 S	M048	1500	3	MgO	S	6		25.44
E0102 S	M048	1600	3	MgO	S	5	0.03(1)	95.90
E110 S	M006	1450	3	MgO	Sn	5	0.3(1)	32.84
E126 S	M006	1450	3	MgO	Sn	5		198.7
E104 S	M047	1300	3	MgO	S+Sn	5	0.5(1)	92.77
E109 S	M047	1400	3	MgO	S+Sn	5	0.02(1)	96.23
E128 S	M048	1300	3	MgO	S+Sn	5		22.47
E134 S	M133	1600	3	MgO	-	5	0.02(1)	9.790

E115 S	M048	1400	3	C	S	5	8.207
E141 S	M048	1400	3	C	S	5	7.015
<i>LA-ICPMS only</i>							
E017	M006+M011		3				0.13(6)
E135	M048		3				0.02(1)
E029	M006+0.01 wt.% S		3				8(7)
E041	M006+1 wt.% S		3				0.16(7)
E041 (sul-sil)	M006+1 wt.% S		3				0.16(7)
E031	M006+1 wt.% S		3				0.3(5)

^a Number of isotopic analysis.

^b Ruthenium concentrations of silicate glasses determined by LA-ICPMS. Uncertainties in parentheses are calculated ... of 5 to 15 spot analysis

^c Ruthenium concentrations determined from the double spike isotope ratio analyses. Uncertainties are <1% including measurement uncertain

^d Measured Ru isotope mass fractionation relative to an Alfa Aesar standard solution. Given uncertainties are 95% c.i. for samples with N≥4.

^e Isotopic difference of metal and nonmetal, which would be depending on the dopant either silicate (no dopant), sulfide (s), a Sn-rich phase (

^f Partition coefficients $D_{Ru}^{metal-silicate}$ calculated using LA-ICPMS concentration data of metals (Table 2) and silicates for selected experiments.

Concentrations were not only determined on the bulk sample (MC-ICPMS), but also in-situ using LA-ICPMS. Partition coefficients (log $D_{Ru}^{metal-silicate}$) were

$\delta^{102/99}Ru^d$	$\Delta^{102/99}Ru_{nonmetal-metal}^e$	$\log D_{Ru}^{metal-silicate}^f$
-----------------------	--	----------------------------------

0.03 ± 0.02

0.04 ± 0.01

0.02 ± 0.02

0.05 ± 0.02

0.07 ± 0.02

0.04 ± 0.02

0.03 ± 0.02

0.05 ± 0.01

0.04 ± 0.02

0.03 ± 0.01

0.03 ± 0.01

0.04 ± 0.02

0.04 ± 0.01

-0.07 ± 0.03

0.11 ± 0.03

-0.03 ± 0.01

0.08 ± 0.01

-0.05 ± 0.05

0.09 ± 0.05^g

0.09 ± 0.03

-0.06 ± 0.01

0.10 ± 0.01

-0.01 ± 0.01

0.05 ± 0.01

0.22 ± 0.01

-0.18 ± 0.01

0.21 ± 0.01

-0.17 ± 0.01

0.05 ± 0.03

-0.01 ± 0.03

-0.01 ± 0.02

0.05 ± 0.02

0.06 ± 0.01

-0.02 ± 0.01

0.02 ± 0.02

0.02 ± 0.02

0.03 ± 0.02 0.01 ± 0.02
0.05 ± 0.01 0.00 ± 0.01

5.9(2)
6.8(3)
2.0(1)
5.6(2)
4.6(2)
5.5(2)

; per sample.

inties and weighing.

{Sn) or a combination of Sn-rich phase and sulfide (S+Sn)

· calculated for samples analyzed by LA-ICPMS only. Capsule C = graphite, t = run duration in hours.