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1	Implications for behavio	or of volatile elements during
2	impacts – zinc and copper	systematics in sediments from
3	the Ries impact structure	and central European tektites
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Abstract

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Moldavites are tektites genetically related to the Ries impact structure, located in Central 29 Europe, but the source materials and the processes related to the chemical fractionation of 30 moldavites are not fully constrained. To further understand moldavite genesis, the Cu and Zn 31 abundances and isotope compositions were measured in a suite of tektites from four different 32 sub-strewn fields (South Bohemia, Moravia, Cheb Basin, Lusatia) and chemically diverse 33 sediments from the surroundings of the Ries impact structure. Moldavites are slightly depleted 34 in Zn (~10–20%) and distinctly depleted in Cu (>90%) relative to supposed sedimentary 35 precursors. Moreover, the moldavites show a wide range in δ^{66} Zn values between 1.7 and 36 3.7‰ (relative to JMC 3-0749 Lyon) and δ^{65} Cu values between 1.6 and 12.5‰ (relative to 37 NIST SRM 976) and are thus enriched in heavy isotopes relative to their possible parent 38 sedimentary sources (δ^{66} Zn = -0.07 to +0.64‰; δ^{65} Cu = -0.4 to +0.7‰). In particular, the 39 Cheb Basin moldavites show some of the highest δ^{65} Cu values (up to 12.5‰) ever observed 40 in natural samples. The relative magnitude of isotope fractionation for Cu and Zn seen here is 41 opposite to oxygen-poor environments such as Moon where Zn is significantly more 42 isotopically fractionated than Cu. One possibility is that monovalent Cu diffuses faster than 43 divalent Zn in the reduced melt and diffusion will not affect extent of Zn isotope 44 fractionation. These observations imply that the capability of forming redox environment may 45 aid in volatilizing some elements, accompanied by isotope fractionation, during the impact 46 process. The larger extent of elemental depletion, coupled with isotope fractionation of more 47 refractory Cu relative to Zn, may also hinge on the presence of carbonyl species of transition 48 metals and electromagnetic charge, which could exist in the impact-induced high-velocity jet 49 of vapor and melts. 50

- *Keywords:* zinc isotopes; copper isotopes; tektites; Ries crater; Ries area sediments; isotope
- 53 fractionation; impact; volatile loss

1. Introduction

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Tektites are terrestrial silica-rich glasses produced during hypervelocity impacts of an 57 extraterrestrial projectile onto the Earth's surface (Koeberl, 1994, 2014; Artemieva, 2008) and 58 as such, they represent a unique source of information about conditions and processes 59 occurring during early phases of the impact process. The close similarity between the 60 chemical composition of the Earth's surface sediments and tektites implies that tektites were 61 formed by reworking of such target materials, mostly from the uppermost unconsolidated 62 strata (e.g., Shaw and Wasserburg, 1982; Koeberl and Fredriksson, 1986; Blum et al., 1992; 63 Dressler and Reimold, 2001; Řanda et al., 2008; Skála et al., 2009; Stöffler et al., 2013; Žák et 64 al., 2016). At present, four major geographically distinct tektite strewn fields are recognized: 65 North American, Central European, Ivory Coast and Australasian. The Central European 66 tektites (moldavites) are assumed to have been formed by reworking of the chemically and 67 68 mineralogically variable sediments of the Upper Freshwater Molasse (Obere Süßwassermolasse, OSM; e.g., Luft, 1983; von Engelhardt et al., 1987, 2005; Žák et al., 69 2016). The Ries impact structure (centered close to Nördlingen, south-eastern Germany) is 70 accepted as the parent crater to moldavites, based on the excellent concordance of their ages 71 at ca. 14.75 ± 0.20 Ma (e.g., Buchner et al., 2010; Schwarz and Lippolt, 2014). However, the 72 agreement between the chemical composition of the OSM sediments and moldavites is not 73 74 excellent, which was explained as a result of chemical modifications during the process of tektite formation (von Engelhardt et al., 2005; Žák et al., 2016). 75 Zinc (lithophile) and Cu (chalcophile) both are volatile/moderately volatile elements 76

77 (Lodders, 2003). Chondrites are significantly enriched in Zn compared with the Bulk Silicate

Earth (BSE) and Mars (McDonough and Sun, 1995; Barrat et al., 2012; Paniello et al., 2012a)

vhile both the Moon and the howardite-eucrite-diogenite (HED) meteorites from asteroid 4-

80	Vesta are depleted in Zn (Paniello et al., 2012a; Paniello et al., 2012b), implying planetary-
81	scale processes leading to sizeable loss of Zn, perhaps caused by giant collisions, thermal
82	metamorphism and/or cataclysmic bombardment (Day and Moynier, 2014). Previously
83	published Zn concentrations in tektites revealed a significant range with a particular
84	difference among individual strewn fields (7–283 ppm; Koeberl, 1992; Koeberl et al., 1997;
85	Moynier et al., 2009a; Skála et al., 2009) and high $\delta^{66/64}$ Zn values from 0.2‰ to 2.5‰ (per
86	mil deviation of the 66 Zn/ 64 Zn ratio from the JMC 3-0749L standard), when compared to the
87	average upper continental crust ($\delta^{66/64}$ Zn from 0‰ to 0.7‰; Albarède, 2004; Chen et al.,
88	2013) and Bulk Silicate Earth ($\delta^{66/64}$ Zn _{BSE} = 0.30 ± 0.03‰; Moynier et al., 2011; Chen et al.,
89	2013; Doucet et al., 2016; Wang et al., 2017). Most tektites analyzed previously show an
90	average Cu content of only 2 ± 1 ppm and are significantly enriched in heavy Cu (δ^{65} Cu
91	values between 2 and 7‰, relative to NIST 976; Moynier et al., 2010) which is in contrast to
92	the results of Li et al. (2009), Liu at el. (2015) and Savage et al. (2015) which, in general,
93	attest to a homogeneous Cu isotope composition of terrestrial igneous lithologies
94	(0.08±0.17‰; Moynier et al., 2017) as well as clastic sediments (0.08±0.20‰; Moynier et al.,
95	2017). Because tektites suffer from minor Zn loss but significant Cu depletion (Moynier et al.,
96	2009a; Moynier et al., 2010; Žák et al., 2016), the impact processes can potentially be used as
97	a proxy for the loss of volatile species during planetary formation.
98	Here, we present new Cu-Zn isotope data for the set of well-characterized moldavites
99	from the Czech Republic and Germany, and chemically diverse sediments from the close
100	surroundings of the Ries impact structure (Žák et al., 2016) in order to provide further
101	constraints on (i) Zn and Cu elemental and isotope variability in possible tektite source
102	materials that could be linked to the heterogeneity of their bulk chemistry, and (ii) the extent
103	of Zn and Cu isotope fractionation between the plausible source sediments and tektites. The
104	latter has been only marginally studied for Zn, with the Ivory Coast strewn field displaying a

105	$\delta^{66/64}$ Zn offset >1‰ between the Bosumtwi crater rocks, having crustal Zn isotope signature,
106	and isotopically heavier Ivory Coast tektites (Moynier et al., 2009a). Moreover, there may be
107	a difference between Bosumtwi and Ries in the quantity of volatiles (in particular water) in
108	the surface layer of the target materials (crystalline versus loosely bound; Koeberl et al., 1998;
109	Žák et al., 2016) which may play a key role in effective fragmentation of molten tektite glass
110	due to the escape of volatile species (see recent model of Žák et al., 2016) and the attainment
111	of conditions suitable for Zn and Cu loss and related isotope fractionation.

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2. Samples and analytical techniques

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In this study, 13 samples of moldavites from four distinct sub-strewn fields of the Central 115 European tektite field and 16 samples of sediments from surrounding of the Ries, 116 117 characterized previously with respect to their major and trace element chemistry, physical properties (shape, color and sculpture) and Li elemental/isotope systematics (Randa et al., 118 2008; Skála et al., 2009; Skála et al., 2010; Řanda et al., 2014; Rodovská et al., 2016; Žák et 119 al., 2016), were analyzed for Zn and Cu abundance and isotope compositions. Details on the 120 general chemistry of moldavites can be found elsewhere (e.g., Bouška et al., 1973; von 121 122 Engelhardt et al., 1987, 2005; Meisel et al., 1997; Trnka and Houzar, 2002; Randa et al., 2008; Skála et al., 2009; Magna et al., 2011; Žák et al., 2016). The studied sediment samples 123 124 were previously characterized with respect to their chemistry and mineralogy (Žák et al., 125 2016). They comprise OSM sediments with variable proportions of fine-grained (clay) fraction and sand fraction, and with variable proportions of quartz and other minerals in the 126 sand fraction (Table 1). Several samples which could have been also present at the surface 127 128 during the impact were also included (sedimentary fill of paleokarst depressions rich in limestone weathering residual minerals; freshwater Oligocene limestone). Sediment sample 129

locations, descriptions, chemical and mineralogical characteristics are given in Žák et al. 130 131 (2016). It should be noted that the exact morphology and sedimentary cover conditions of the target area itself are unknown, because they were completely destroyed by the impact. 132 Therefore, the studied sediments represent only an approximation of the conditions and 133 materials of the intrinsic target area. In addition, three samples of clay-size fractions, isolated 134 from sedimentary samples using a common sedimentation method in distilled water were 135 136 measured (we note that these fractions are not completely pure clay fractions but may contain a small proportion of other minerals including secondary phases, such as secondary Fe and 137 Mn oxides/hydroxides). The procedure to prepare samples for clay minerals identification 138 139 involved extensive washing in distilled water and separation of the clay fraction in the Andreasen cylinder. To determine bulk mineral composition X-ray diffraction patterns have 140 been acquired from untreated bulk specimens between 3 and 70° 2 Θ (CuK α_1 radiation). The 141 specimens have been scanned with a diffractometer between 2 and $40^{\circ} 2\Theta$ and then saturated 142 with ethylenglycol for 4 hours at 80°C. Following saturation the new set of X-ray diffraction 143 data has been acquired. After that, the glass sample wafers with specimens have been heated 144 at 550°C under ambient atmosphere for 1 hour. In general, clay mineral composition is similar 145 in all samples with subtle differences among individual sediments. Identified clay minerals 146 147 include kaolinite, illite, chlorite, montmorillonite and interstratified illite/montmorillonite. The analytical procedures for Zn and Cu purification were performed in the CAGE 148 149 laboratories, the Institut de Physique du Globe de Paris (IPGP). Zinc was isolated and purified by a two-stage HBr-based anion exchange chromatography following methods given 150 151 elsewhere (Moynier et al., 2006; Moynier and Le Borgne, 2015). A single-stage anion exchange chromatography for Cu purification using AG-MP1 resin was employed (Maréchal 152 153 et al., 1999; Savage et al., 2015) with 95–100% recovery; the total procedural blank for Cu was ~4 ng. The Zn and Cu concentrations and isotope compositions were measured by 154

155	standard-sample bracketing (Maréchal et al., 1999; Moynier et al., 2006, 2009a, 2010; Savage
156	et al., 2015) using a Neptune Plus multiple-collector inductively-coupled-plasma mass
157	spectrometer (MC-ICPMS; ThermoFisher, Bremen, Germany), housed at IPGP. The 2SD on
158	repeated measurements of the bracketing standard was $\pm 0.06\%$; for multiple dissolutions and
159	purifications of BHVO-2, the 2SD uncertainty was $\pm 0.08\%$. Zinc and Cu concentrations were
160	determined against a 100 ppb Zn reference solution (JMC 3-0749 Lyon) and a 50 ppb NIST
161	SRM 976 reference solution, respectively. Zinc isotope compositions were calculated as δ^{x} Zn
162	$(\%) = [(^{\times}Zn/^{64}Zn)_{sample}/(^{\times}Zn/^{64}Zn)_{standard} -1] \times 1000$, where $^{\times}Zn$ denotes ^{66}Zn , ^{67}Zn and ^{68}Zn ,
163	respectively. Potential interferences from ²³ Na ⁴⁰ Ar ⁺ and ²⁵ Mg ⁴⁰ Ar ⁺ on ⁶³ Cu and ⁶⁵ Cu,
164	respectively, were ruled out by measuring Na/Cu and Mg/Cu ratios in the samples prior to the
165	isotope analysis. Isotope ratios of Cu are expressed as δ^{65} Cu (‰) =
166	$[(^{65}Cu/^{63}Cu)_{sample}/(^{65}Cu/^{63}Cu)_{standard} -1] \times 1000$. Reference rock standards BCR-2 and BHVO-
167	2 (US Geol. Surv.) were processed and analyzed along with the unknown samples to monitor
168	the accuracy and reproducibility of the analytical procedures and spectrometry routines; their
169	Zn and Cu isotope compositions were consistent with previous studies (Table 1 and Archer
170	and Vance, 2004; Moynier et al., 2010; Moeller et al., 2012; Chen et al., 2013; Sossi et al.,
171	2014; Savage et al., 2015; Lv et al., 2016).
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173	3. Results
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175	3.1 Zinc abundance and isotope compositions
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177	The Zn concentrations and isotope ratios in moldavites, sediments and clay fractions are
178	reported in Table 1 and plotted in Fig. 1. The moldavites show a wide range in Zn abundance
179	(1.9–108 ppm), extending to both lower and higher Zn contents compared with data from

Moynier et al. (2009a), which may reflect the wider chemical range of moldavites selected for 180 this study (see Skála et al., 2009). Moldavites from South Bohemia have 1.9-34.5 ppm Zn, 181 those from Moravia have 7.3–18.8 ppm Zn and those from the Cheb Basin contain 20.6–108 182 ppm Zn (with three out of four samples having [Zn] > 58 ppm); a single moldavite from 183 Lusatia has 16.7 ppm Zn. These results attest to modest differences between the Cheb Basin 184 moldavites and those from the other sub-strewn fields, noted by Řanda et al. (2008) and Skála 185 et al. (2009). The Ries sediments show a remarkable range in Zn abundances from 4.6 to 186 186 ppm Zn. Zinc contents in sediments correlate with Al₂O₃ contents (Fig. 2) and FeO (not 187 shown). Zinc contents obtained in this study are broadly consistent with those reported by Žák 188 189 et al. (2016) for different aliquots of the same samples of moldavites and sediments, with the exception of slightly lower Zn contents in high-Zn samples reported in this study. The clay 190 fractions from three sediments (1 UTZ, 10 GLM, 13 BMB) have from 94 to 161 ppm Zn; this 191 192 is, in all three cases, higher than Zn abundance in the respective bulk sample, in case of sample 13 BMB by a factor of three (Table 1). 193

The moldavites in this study show a range in δ^{66} Zn = 1.7–3.7‰, which is significantly 194 larger compared to data reported by Moynier et al. (2009a) for four south Bohemian 195 specimens (δ^{66} Zn = 1.9–2.3‰). The moldavites from different sub-strewn fields do not show 196 distinct Zn isotope compositions (Moravia: δ^{66} Zn = 2.1–2.7‰; Lusatia: δ^{66} Zn = 2.5‰; South 197 Bohemia: δ^{66} Zn = 1.7–3.3‰; the Cheb Basin: δ^{66} Zn = 2.0–3.7‰). The sediments from the 198 Ries surroundings show little variability in δ^{66} Zn values (-0.07‰ to 0.64‰), which is 199 consistent with global data for sediments (see recent review byMoynier et al., 2017). The clay 200 fractions always have lighter Zn isotope compositions than the corresponding bulk sediments, 201 and define a narrow range of Δ^{66} Zn_{clav-bulk sediment} values (δ^{66} Zn_{clav} – δ^{66} Zn_{bulk sediment}) between 202 203 -0.5 and -0.7%, indicating preferential incorporation of light Zn isotopes into clay.

3.2 Copper abundance and isotope compositions

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The Cu contents and isotope compositions are listed in Table 1 and plotted in Fig. 1. The 207 moldavites show a range in Cu abundance from 0.2 to 2.9 ppm (the values were taken from 208 Žák et al., 2016), much lower than the canonical abundance of the upper continental crust at 209 ~28 ppm (Rudnick and Gao, 2014). Moldavites from South Bohemia have 0.2–1.4 ppm Cu, 210 those from Moravia have 0.6–0.8 ppm Cu and those from the Cheb Basin have 0.4–2.9 ppm 211 Cu; a single Lusatian moldavite has 1.1 ppm Cu. These values are consistent with those found 212 by Skála et al. (2009) for the Cheb Basin moldavites (0.9 to 4.7 ppm) as well as for other 213 214 tektite fields, with the exception of low Cu contents for moldavites (<0.4 ppm) reported by Moynier et al. (2010). The Ries sediments show a range in Cu concentrations from 0.8 to 58 215 ppm Cu; these values are consistent with those reported by Žák et al. (2016). The clay 216 217 fractions from three sediments have from 21.4 to 42.2 ppm Cu which is in all three cases higher than Cu content in the respective bulk sediment, and is consistent with the findings for 218 Zn. 219 The moldavites from South Bohemia have δ^{65} Cu values from 2.2 to 5.4‰, those from 220 Moravia between 5.8 and 8.8‰ and those from the Cheb Basin between 1.6 and 12.5‰; a 221 single Lusatian moldavite has δ^{65} Cu of 6.1‰. The new δ^{65} Cu values measured for moldavites 222

extend towards significantly heavier values than those reported previously (Moynier et al.,

224 2010), with the Cheb Basin moldavites showing some of the highest δ^{65} Cu values (up to

12.5‰) ever observed in natural high-temperature modified samples. The largest previously

reported natural ⁶⁵Cu enrichments were observed in strongly weathered shales from central

227 China (up to 19.7%; Lv et al., 2016) and secondary Cu minerals associated with ore body

formation, where δ^{65} Cu values are typically positive in the supergene, and can reach ~10%

(e.g., Rouxel et al., 2004; Mathur et al., 2009). The δ^{65} Cu values in the Ries area sediments

230	show a narrow variation (-0.4 to 0.7‰) and are in the range of common crustal lithologies
231	(Luck et al., 2003; Li et al., 2009). The clay fractions show a range in δ^{65} Cu values between
232	-0.1 and $+0.2$ % and are, contrary to Zn, isotopically heavier than the respective bulk
233	sediments (Δ^{65} Cu _{clay-bulk sediment} values between 0.01 and 0.21‰).
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235	4. Discussion
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237	4.1.1 Statistical evaluation of Zn isotope data – implications for isotope fractionation in
238	natural processes
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240	It has been discussed that kinetic effects may become an important factor in forming
241	measurable isotope fractionation effects. For example, (Young and Galy, 2004) raised this
242	possibility for Mg isotopes although the resolution of equilibrium and kinetic fractionation
243	processes in the three-isotope Mg plot seems to be barely outside of the statistical resolution.
244	Applying the approach of (Young and Galy, 2004) to Zn, the calculated theoretical slopes β =
245	1.942 and 1.972 for equilibrium and kinetic fractionation, respectively, suggest that these two
246	different mechanisms of Zn isotope fractionation could be resolved in nature. With the given
247	data set we were able to test this hypothesis for Zn isotopes. In the triple-Zn isotope plot (Fig.
248	3), both <i>x</i> - <i>y</i> variables (i.e., δ^{66} Zn and δ^{68} Zn) can be considered as random, containing
249	measurement errors and other variability. Such cases are referred to as structural or model II
250	regression. A very simple structural relationship is assumed to be hidden in our data in the
251	form of a line with theoretically known slope. The SMATR package (Falster et al., 2006;
252	Warton et al., 2006) provides the possibility of fitting bivariate lines using the major axis
253	(MA) or standardized major axis (SMA).

In the present study, we defined three separate groups, sediments (S), moldavites from 254 255 this study (M), and other published tektites (T). The results of fitting MA lines, paralleled by 95% confidence intervals, are listed in Table 2. Because the basic assumption for MA 256 regression is the bivariate normality, we have inspected whether the S, M and T group data 257 are approximately bivariate normal, and applied the normality tests. For the group M, the 258 normality assumption was violated and two samples SBM-192 and SBM-44 were excluded 259 from further statistical evaluation. The former sample represents only a single measurement 260 and averaging of several independent measurements was therefore not possible, while the 261 latter is an average of three measurements all of which deviated outside the pattern of the 262 263 individual measurements of the other samples in this study. The cleaned data were denoted as M1 (SBM-192 excluded) and M2 (both SBM-192 and SBM-44 excluded), and the results for 264 all the three variants of moldavite data (M, M1, and M2) are listed in Table 2. Furthermore, a 265 bootstrap (by resampling residuals) was applied to compute confidence intervals for the 266 slopes and elevations for comparison. A good agreement of the computed slopes b with the 267 theoretical ones ($\beta = 1.942$ for sediments, and 1.972 for moldavites and other tektites), is 268 observed with all confidence intervals containing the theoretical values. In case of groups M1 269 270 and M2 the confidence intervals are narrower. The computed intercepts a are close to zero with a larger difference for the group M. 271

As the lines for all groups should pass through the origin, we have also used MA regression with zero intercept (i.e., with the line forced through the origin). For the group M the theoretical slope (1.972) is not in the 95% confidence interval, implying statistically significant difference. Due to violation of the normality assumption for the group M, the cleaned data M1 and M2 reveal an agreement with the theoretical slope. For the group T, the regression with zero intercept shows a significant difference from the theoretical slope. The reasons for this difference are unknown but some inter-laboratory differences could perhaps

be involved. We tested several inter-group comparison hypotheses using SMATR. (i) By 279 280 comparing S and M (M1, M2) groups only (i.e., data from this study measured in the same laboratory), the test for common slope did not reject the hypothesis that slopes are equal (p =281 0.082, 0.226, 0.384). (ii) The test for particular slope did not reject the hypothesis that the 282 common slope of other tektites (T) and moldavites M (M1, M2) is equal to 1.97 (p = 0.071, p = 0.071)283 0.141, 0.165). (iii) The test for common slope did not reject the hypothesis that slopes of all S, 284 M (M1, M2), and T are equal (p = 0.066, 0.092, 0.085). Collectively, the detailed statistical 285 treatment indicates that a significant statistical difference between sediments and tektites was 286 not proven (Fig. 3) but in some cases, this hypothesis was close to rejection (Table 2). Such a 287 288 conclusion leaves the question of possible resolution of Zn isotope fractionation during kinetic and equilibrium processes unsolved. It may be that this subtle difference may only 289 become apparent at Zn isotope compositions that are much beyond the range of common 290 291 terrestrial as well extra-terrestrial values.

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4.2 Zinc and copper systematics in the Ries area sediments

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The δ^{66} Zn variation in common terrestrial crustal igneous rocks is, in general, limited to 295 between ~0 and 0.6‰ (Maréchal et al., 1999; Chen et al., 2013), and can be up to 0.7‰ in 296 sedimentary rocks (Pichat et al., 2003; Albarède, 2004; Mason et al., 2005). This is paralleled 297 by moderately high Zn abundance in the upper continental crust (67 ± 6 ppm; Rudnick and 298 Gao, 2014) compared with ~55 ppm in the primitive upper mantle (McDonough and Sun, 299 1995), and between ~20 and 200 ppm in different granite types (I, S, A; (Hall, 1975; Chappell 300 and White, 1992; King et al., 1997). These findings are consistent with uniform Zn isotope 301 composition of the bulk Ries sediments in this study (Fig. 1) with δ^{66} Zn constrained between 302 -0.07 and 0.64‰. On the other hand, Zn contents in sediments from the Ries vary 303

significantly by a factor of ~40 which, together with the variability of Zn contents of 304 305 moldavites, attests to a compositional variability of the target materials and incorporation of chemically distinct sedimentary lithologies that ultimately led to the formation of moldavites 306 (e.g., von Engelhardt et al., 1987; Rodovská et al., 2016; Žák et al., 2016). The apparently 307 higher Zn contents in the Cheb Basin moldavites suggest that specific sediments must have 308 been sourced which also have higher Fe and Al abundances (Fig. 2). The paleogeography 309 310 indicates that the northern boundary of the OSM sediments occurred in the target area (e.g., Pohl et al., 1977; Hörz et al., 1983). Uneven karst surface with residual karst sediments could 311 have been present in the northern part of the target area. This type of material, represented by 312 the samples 14 BMB, 24 RTS and 25 RTS, shows distinctly higher Zn contents (>118 ppm), 313 paralleled by elevated abundances of Cu, Al, and Fe. Their participation in the ultimate source 314 of materials for moldavites of the Cheb Basin area, carrying elevated Zn contents, is therefore 315 316 probable (see also Rodovská et al., 2016).

Typical Cu contents in continental crustal lithologies vary greatly (~28 ppm in the bulk 317 continental crust, ~118 ppm in shales, ~20 ppm in soils; Jeng, 1992; McDonough and Sun, 318 1995; Bigalke et al., 2011; Rudnick and Gao, 2014) while ultramafic mantle rocks have 20-30 319 ppm Cu (McDonough and Sun, 1995; Palme and O'Neill, 2014). The effusive and intrusive 320 321 igneous rocks from the Earth's mantle and crust, such as basalts and granites, typically have invariant δ^{65} Cu values of ~0‰ (Luck et al., 2003; Archer and Vance, 2004; Li et al., 2009; 322 Savage et al., 2015), and the weighted mean δ^{65} Cu of continental sediments of ~0‰ attests to 323 insignificant Cu isotope fractionation during weathering (note a restricted range in δ^{65} Cu 324 325 between -0.57‰ and 0.44‰ found for different soil horizons; Bigalke et al., 2011). In contrast, recent data of Lv et al. (2016) reveals extreme fractionation of Cu isotopes during 326 chemical weathering, with δ^{65} Cu values ranging between -6.4 and 19.7‰. This is 327 underscored by significant Cu isotope fractionations reported for oxidative weathering of 328

primary sulphides (0.0 to $\pm 2.0\%$ δ^{65} Cu_{solution-chalcopyrite rock}; Fernandez and Borrok, 2009; 329 Mathur et al., 2012), as well as during sorption on Fe-oxides and organic complexation 330 331 (Fernandez and Borrok, 2009). Lithologically diverse sediments in this study show a range in Cu concentrations between 0.9 and 58 ppm (Table 1) that are well correlated with Zn 332 333 contents (only the sample 15 HRD shows elevated Cu abundance at a given Zn content; Fig. 334 1a). The broad Zn–Cu isotope co-variation observed for sediments from surroundings of the 335 Ries (Fig. 1b) also indicates a similar sense of stable isotope fractionation of chalcophile elements during weathering of common silicate lithologies despite the possibility that Zn and 336 337 Cu might be accommodated in different phases in the source rocks of sediments and in the secondary minerals. The contents of Cu in sediments correlate with FeO, Al₂O₃ and suggest 338 higher FeO and Al₂O₃ content in possible parent sediments. This would indicate somewhat 339 different behavior of Zn and Cu during weathering in the sense that these elements would be 340 isotopically fractionated in the opposite direction during formation of clay minerals. This 341 342 means that (i) the formation of unconsolidated materials through weathering imparts rather minor isotope effects to both Zn and Cu, (ii) the behavior of Cu and Zn during tektite 343 formation is not completely identical, and (iii) strong isotope effects observed for tektites (see 344 345 Section 4.3) must solely be linked to conditions existing during or shortly after the impact. 346

347 4.2.1 Behavior of Zn and Cu at low temperatures

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Primary and secondary mineralogy can play a determining role in the stable isotope systematics of sedimentary units although for many isotope systems the experimental data and/or natural observations are insufficient or absent. Clay fractions isolated from bulk sediments are uniformly enriched in Cu and Zn relative to the corresponding bulk sediments despite the compositional diversity (clay-rich, quartz sand-rich, limestone weathering

354	residuum), attesting to similar behaviour of Cu and Zn at low temperatures, although
355	adsorption of heavy metals may differ upon variable chemical and physical conditions (e.g.,
356	Young Do and Park, 2011; Helios-Rybicka and Wójcik, 2012). The enrichment of Cu and Zn
357	is most apparent for the low-[Cu]–[Zn] sample 13 BMB with an enrichment factor in the clay
358	fraction of >2 and ~3 for Cu and Zn, respectively. Notable differences are, however, observed
359	for Zn and Cu isotope systematics in clay fractions. Zinc isotope compositions are
360	significantly lighter in clays relative to the corresponding bulk sediments which is in contrast
361	to the findings of Guinoiseau et al. (2016) who observed a systematic enrichment in heavy Zn
362	isotopes on the surface of kaolinite (Δ^{66} Zn _{adsorbed-solution} from 0.11‰ to 0.49‰).
363	Cu isotopes in clay fractions show zero to slight enrichments in ⁶⁵ Cu. This indicates a
364	mineralogy-mediated dichotomy in generating intra-sample Zn–Cu isotope variations, which
365	may perhaps reflect occupation at different crystal sites for Zn and Cu. Still, δ^{66} Zn values in
366	clay fractions are similar to those reported for global clays (e.g., Maréchal et al., 2000).
367	Interestingly, the recent experimental study of Li et al. (2015) showed the opposite sense of
368	Cu isotope fractionation during adsorption onto kaolinite where the Δ^{65} Cu _{kaolinite-solution} of
369	$-0.17 \pm 0.10\%$ (2 σ) appears to invoke preferential incorporation of isotopically light Cu into
370	clays, perhaps as a result of sorption of free Cu ion onto the surface of clay mineral. Li et al.
371	(2015) concluded that the isotope compositions of Cu adsorbed onto natural soils may vary
372	greatly at relatively low Cu concentrations of the soil solutions. This is underscored by a large
373	variation in δ^{65} Cu from -17 to +10‰ that was observed in secondary mineral phases which
374	typically develop during weathering of sulphides in the near-surface environment (Mathur et
375	al., 2009). These observations indicate that clay minerals of the OSM sediments cannot form a
376	major part of the moldavite melts because they are distinctly enriched in Zn, and, at the same
377	time, they carry a resolvedly light Zn isotope signature (Fig. 1). Moreover, they also have too
378	much Al and not enough Si so that their theoretically elevated proportion would violate major

element constraints because only the sands (containing abundant mica, feldspar, etc.) with a
moderate clay admixture and a very small content of clastic (mainly dolomitic) carbonates are
suitable for moldavite production (von Engelhardt et al., 1987; Žák et al., 2016).

382

4.3 Zinc and copper systematics in moldavites – implications for the impact processes

384

Several stable isotope systems have been studied previously (Humayun and Clayton, 385 1995; Wombacher et al., 2003; Moynier et al., 2009a, 2010; Magna et al., 2011; Rodovská et 386 al., 2016 in order to test the possible isotope fractionations related to extreme physical 387 388 conditions existing during the hypervelocity impact events. For example, Moynier et al. (2009a) have found large variations for Zn isotopes (δ^{66} Zn = 0.3–2.5‰), and the differences 389 in δ^{66} Zn among moldavites, Ivory Coast tektites, Australasian tektites (including Muong 390 391 Nong-type tektites), and bediasites (North American tektite field) were explained as a result of evaporation in the context of shock heating. However, the p-T conditions of individual 392 impacts and the pre-impact chemical and lithological situation may be largely different, which 393 could perhaps be reflected in variable Zn isotope systematics of tektites from individual 394 strewn fields. In a follow-up study, Moynier et al. (2010) explored Cu elemental and isotope 395 systematics in tektites and have shown that (i) Cu in tektites was significantly more depleted 396 than Zn, and (ii) tektites displayed a significantly larger range in δ^{65} Cu than in δ^{66} Zn 397 (reflected in larger ‰/amu variation) despite higher half mass condensation temperature of 398 Cu compared to Zn. Indeed, Albarede et al. (2007) noted that Cu is isotopically more 399 fractionated (up to 0.8‰) than Zn (up to 0.4‰) in shocked sandstones from the Meteor 400 Crater, USA. This could, in part, result from the differences in ionic radii of Zn²⁺ and Cu⁺ 401 which translates into different diffusion coefficients of these ions in silicate melts (see 402 discussion in Moynier et al., 2010). Highly reducing conditions of the impact systems have 403

been documented by the predominance of Fe^{2+} over Fe^{3+} in moldavites (e.g., Dunlap et al., 404 1998; Skála et al., 2009) or reduced gaseous species (Žák et al., 2012), while in the source 405 sediments this relationship is opposite; the existence of Cu^+ rather than Cu^{2+} compositions is 406 therefore likely during tektite-forming events. Wombacher et al. (2003) have found significant 407 enrichments in heavy Cd isotopes in one investigated Muong Nong-type tektite caused 408 probably by evaporative loss whereas other terrestrial materials have shown no or very minor 409 variation in $\delta^{112/114}$ Cd. Combined studies of Li systematics in central European tektites and 410 possible target sediments from the Ries (Magna et al., 2011; Rodovská et al., 2016) indicated 411 412 that neither is Li lost during the impact event nor are Li isotopes fractionated significantly by shock heating and flash melting. Moreover, it appears that Li is homogenized effectively in 413 tektites despite a large range in Li contents and isotope compositions of target sediments. 414 Studies of somewhat more volatile potassium (Humayun and Clayton, 1995; Herzog et al., 415 2008) are consistent with a limited loss of some elements from a molten glass into the 416 417 vapor/plasma although energetic collisions may fractionate K isotopes (Wang and Jacobsen, 2016). 418

In contrast to sediments (Section 4.2), all tektites measured in the present study are 419 distinctly enriched in the heavy isotopes of Zn and Cu (Fig. 1b), consistent with earlier results 420 (Moynier et al., 2009a, 2010). The only other natural samples known so far to have 421 comparably heavy Zn isotope compositions are lunar regolith samples (δ^{66} Zn up to 6.4%); 422 Moynier et al., 2006; Paniello et al., 2012a; Kato et al., 2015), EL6 enstatite chondrites and 423 eucrites (Moynier et al., 2011; Paniello et al., 2012b) although not all of these high- δ^{66} Zn 424 samples suffered from a resolvable Zn loss. In particular, Hvittis, Atlanta and North West 425 Forrest EL6 enstatite chondrites have δ^{66} Zn values up to 7.4‰, the highest observed so far for 426 Solar System materials, with very low Zn concentrations compared to enstatite chondrites of 427 lower metamorphic grades (Moynier et al. 2011). Recent studies have shown that Zn isotopes 428

are significantly fractionated during evaporation-condensation processes (Moynier et al., 429 2011). For example, the δ^{66} Zn of lunar basalts is ~1% heavier than average terrestrial basalts 430 and this extent of fractionation is assumed to have occurred during an evaporation event 431 (Paniello et al., 2012a; Day and Moynier, 2014; Kato et al., 2015). A recent study of Day et 432 al. (2017) has shown that Zn isotopes are indeed fractionated by ~ 0.5 %/a.m.u. at high 433 temperatures and pressures (locally >8400K and >8 GPa) by evaporation during the nuclear 434 435 deronation at the Trinity test site, USA, following a Rayleigh distillation processes and that Zn loss could be significant under some conditions. On the other hand, Zn loss during tektite 436 formation does not exceed ~20% (this study and Žák et al., 2016) and our Zn (and Cu) 437 elemental and isotope data do not fit a simple Rayleigh fractionation process suggesting that 438 evaporation processes taking place during formation of tektites are more complex than during 439 the nuclear detonation. 440

It has been recognized that most chondrite groups are enriched in ⁶³Cu relative to BSE, 441 with δ^{65} Cu values varying from -1.5 to -0.1‰ in carbonaceous chondrites, -0.5 to +0.1‰ in 442 ordinary chondrites and -0.4 to -0.15‰ in enstatite chondrites (Luck et al., 2003; Savage et 443 al., 2015). This observation implies that impact events and accretion processes result in 444 distinct Cu isotope systematics of distal impact melt ejecta and planetesimals. Furthermore, 445 the new δ^{65} Cu values for tektites in this study (up to 12.5‰) are, to our knowledge, some of 446 the highest ever measured in natural samples. These high values cannot originate from a 447 common natural process; for example, smelting is unable to fractionate Cu isotopes despite 448 high temperatures applied (Gale et al., 1999; Mattielli et al., 2006). While the new Zn and Cu 449 data for tektites may appear to support the evaporation-induced fractionation process (see 450 Moynier et al., 2009a, 2010), the lack of possible parental materials to tektites in these studies 451 did not enable a more exact assessment of alternative processes. The full complement of new 452 data for sediments from the Ries (**Table 1**) appears to put certain ambiguity on a purely 453

evaporative loss of chalcophile elements. This is illustrated in Fig. 4 where Rayleigh 454 455 fractionation paths are plotted for several starting Zn and Cu concentrations in order to mimic the large range in Zn and Cu contents of the sedimentary parentage. For Zn, all calculated 456 paths (elemental and binary species) cross-cut the field of moldavite data at a too 457 perpendicular angle, invoking the inability of the Rayleigh process to result in the observed 458 elemental and isotope systematics of Zn. The model also indicates that a singular initial Zn 459 abundance value may be at odds with natural systems. On the other hand, the Cu data can be 460 broadly anchored by a Rayleigh fractionation process if the starting Cu concentration is ~3–4 461 ppm. Such a low Cu content is not excluded from the current data set (Table 1) but we stress 462 463 that our data set only represents the best available proxy for Cu systematics of the sedimentary parentage to moldavites and that moldavite melt represents a mixture of a 464 number of chemically diverse sediments. 465

466

467 **4.4 Implications for volatile loss during textite formation**

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The major observations of this study are two-fold: (i) Zn and Cu are lost to distinctively 469 different degrees from the assumed sedimentary materials with Cu loss being significantly 470 471 more pronounced (up to 95%) than Zn loss ($\sim 10-20\%$), calculated from the average Cu and Zn contents of volatile loss-corrected OSM sediments (von Engelhardt et al., 2005; Žák et al., 472 2016), and (ii) Cu isotopes in tektites are significantly more fractionated than Zn isotopes. 473 474 These findings are consistent with earlier observations made for tektites (Moynier et al., 2009a, 2010) but are opposite to what has been found for lunar samples where Zn is 475 significantly more fractionated than Cu (Moynier et al., 2006). In the model of Moynier et al. 476 (2009a), a velocity field stirs the tektite, while differential evaporation of Zn isotopes occurs 477 at the surface. If the characteristic evaporation time is slow with regard to the diffusion 478

timescale, the system deviates from Rayleigh process because a diffusion-limited regime is established. As a consequence, the magnitude of the Zn isotope fractionation is reduced for the latter process at a given extent of Zn loss. The isotope fractionation in a diffusion-limited regime is controlled by the balance between the evaporative flux and the diffusive flux at the diffusion boundary layer.

The role of redox conditions has been suggested by Moynier et al. (2010) and recent 484 experimental study by (Ni and Zhang, 2016) provides evidence that monovalent Cu is 485 significantly more mobile than divalent Zn in silicate melts, most likely as a consequence of 486 higher chemical diffusivity. This would perhaps imply that oxidation state and chemical 487 488 makeup of the system prior to impact (which would control the mineralogical host and chemical exchange in the impact plasma) may play a prominent role in fractionating these 489 elements during the impacts. In fact, the volatility of the elements is directly related to the 490 491 redox conditions (e.g., O'Neill and Palme, 2008). In the gas phase, both Cu and Zn will be present as mono-atomic Cu_g and Zn_g at a large range of physical conditions (Moynier et al., 492 2007). On the other hand, in the condensed phase Cu will be present as Cu⁺ and, therefore, 493 associated to either 1/2 S atom or 1/2 O atom as CuS_{1/2} or CuO_{1/2} while Zn will be present at 494 Zn²⁺ and associated with one O atom as ZnO in silicate minerals. Following a similar 495 496 approach as applied by O'Neill and Palme (2008), this would indicate that Cu will become relatively more volatile than Zn at higher f_{O_2} and would be consistent with observations from 497 this study. Moreover, significant redox-related diffusive loss of Cu from tektites can be 498 compared with conservative behaviour of Li which belongs to the fastest diffusing elements 499 500 with associated isotope effects (e.g., Richter et al., 2003) and yet, tektites do not show any kinetic effects on Li (Magna et al., 2011; Rodovská et al., 2016). 501 In earlier studies, von Engelhardt et al. (1987, 2005) proposed that moldavites are early 502

503 condensates from the plasma of vaporized source materials in which the elements with larger

ions were preferentially trapped in the condensed silicate melt relative to those with small 504 505 ionic radii. However, this model of vaporization contradicts the presence of partly unmelted inclusions (baddeleyite) and abundant domains of pure silica glass (lechatelierite in 506 moldavites; Trnka and Houzar, 2002). Konta and Mráz (1975) experimentally stepwise 507 melted a sample of sandy-silty clay and suggested that evaporation of the silicate material 508 caused a significant change in relations between the major oxides compared to the original 509 510 chemical-physical relations in the source rocks. They also concluded that direct melting of source sediments is unlikely to form tektites and that evaporation-condensation process 511 should be taken into account. This would require a significant part of the tektite material to 512 513 have been converted to plasma and re-condensed during rapid cooling. The recent model of moldavite formation (Žák et al., 2016) is based on fragmentation of tektite melt due to 514 separation of volatiles (i.e., H₂O, CO₂, CO) during adiabatic decompression of the ejected 515 516 matter, followed by coalescence of molten glass droplets to larger tektite bodies. The temperature, at which the individual moldavite bodies were formed, was in the range between 517 the melting temperature of moldavite glass and the melting temperature of lechatelierite, 518 roughly between ~1100 and ~1713°C. However, this temperature range is too high for both 519 Zn and Cu to condense. Therefore, the process of element loss, instead of condensation, 520 521 should perhaps be considered for elements with T_c lower than that of potassium or rubidium, both of which would unlikely to be condensed on a hot glass. 522

In addition to these observations, we offer further additional physical factors that may help elucidate the Cu–Zn elemental and isotope systematics during the impacts. We note, however, that no experimental data exist for relevant impact and post-impact conditions. It could be that electromagnetic properties of ejected particles may come into play. Monovalent and divalent Cu differ in magnetic properties whereby Cu(I) is diamagnetic and Cu(II) is paramagnetic. Electric charges are generated in particle clouds (Pähtz et al., 2010) but it is not

clear at present, whether such an effect may impart separation of Cu isotopes in the transient 529 530 plasma/overheated melt stage of the matter. Furthermore, Fe-group elements can readily form carbonyls at extreme conditions which are relevant to tektite formation (Řanda et al., 2008). 531 Organic matter was present at the impact site and formation of these highly volatile 532 organometallic compounds cannot be excluded at some stages of the impact process and 533 subsequent p–T relaxation (Cody et al., 2000). Žák et al. (2016) noted that Fe and Cu belong 534 to the few elements that display a large extent of depletion in moldavites relative to possible 535 precursor materials and it remains to be tested whether organometallic compounds may 536 fractionate isotope systems of some other transitional metals to a resolved degree of 537 538 confidence. Preliminary experimental data of Wiesli et al. (2007) have hinted at marginal Fe isotope fractionation of liquid and vapor Fe-pentacarbonyl. However, only further 539 experimental work can provide new limitations to the behavior of stable isotope systems at 540 541 extreme conditions. 542 **5.** Conclusions 543 544 The moldavites measured in this study are depleted in Zn and Cu contents, and are 545 distinctly enriched in the heavy Zn and Cu isotopes relative to their possible parent 546 sedimentary sources with Cu isotope fractionation in tektites being significantly more 547 pronounced than that of Zn. The origin of this is unclear at present but redox transition of 548 divalent to monovalent Cu appears to be the dominant factor, resulting in strikingly different 549 diffusivities of Cu^+ and Zn^{2+} in tektite melts. It could also be potentially related to formation 550 of volatile carbonyls, or perhaps to different electromagnetic properties of monovalent and 551

divalent Cu. Clay fractions isolated from bulk sediments are uniformly enriched in Cu and Zn

relative to the corresponding bulk sediments despite the compositional diversity of the latter.

The clay minerals of the Ries sediments cannot form a major part of the moldavite melts 554 555 because they are enriched in light Zn isotopes. The karst sediments show distinctly higher Zn and Cu contents paralleled by elevated abundances of Al and Fe, and cannot thus be a major 556 lithological composition for moldavites, perhaps with the exception of the Cheb Basin 557 moldavites, where participation of karst-like residual soils in the ultimate source of 558 moldavites is probable. Rayleigh distillation process appears to be incapable of generating the 559 observed Zn and Cu isotope variability in tektites although no singular starting composition 560 should be applied, considering a possible range in chemistry of parent sediments of tektites. 561 From the Zn triple-isotope plot it follows that kinetic effects may become important for high-562 563 temperature processed matter while equilibrium processes apply to low-temperature samples, although sizeable difference could only become apparent at high values of δ^{66} Zn and δ^{68} Zn. 564 The opposite sense of magnitudes of Zn and Cu isotope fractionation in tektites versus lunar 565 samples invokes an important role of oxygen and redox processes in general in inducing 566 567 isotope fractionation of some chalcophile elements whereas other processes apply for oxygendevoid environments. 568

569

570

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			δ ⁶⁶ Zn		δ ⁶⁷ Zn		δ ⁶⁸ Zn		Cu	δ ⁶⁵ Cu		Al ₂ O ₃
	Petrographic description	Zn (ppm)	(‰)	2SD	(‰)	2SD	(‰)	2SD	(ppm)	(‰)	2SD	(wt.%) ^a
Ries area												
1 UTZ	Unterzell; OSM, clay-rich ("bentonite clay")	81.8	0.31	0.08	0.40	0.36	0.57	0.11	17.4	0.00	0.03	13.8
	clay fraction	93.9	-0.52	0.08	-0.74	0.07	-1.01	0.16	26.9	0.21	0.01	-
2 UTZ	Unterzell; OSM, clay- and carbonate-rich	37.9	0.09	0.10	0.24	0.10	0.22	0.12	7.5	0.29	0.04	9.34
3 UTZ	Unterzell; OSM; quartz-dominated fine-grained sand	19.2	0.28	0.13	0.37	0.32	0.47	0.21	3.2	-0.33	0.02	5.07
4 UTZ	Unterzell; OSM, silty-sandy layer	42.0	0.11	0.07	0.26	0.12	0.25	0.14	5.5	-0.11	0.07	9.72
5 ZMTH	Ziemetshausen; OSM, sand with carbonate	18.2	0.17	0.16	0.21	0.34	0.33	0.25	2.1	0.11	0.03	5.79
9 GLM	Guldesmühle; OMM, carbonate and clay-rich sand Guldesmühle; OSM, fine-grained sand with clay and	57.5	0.04	0.03	0.14	0.06	0.11	0.08	2.7	-0.15	0.02	7.79
10 GLM	carbonate	108	0.46	0.12	0.61	0.33	0.86	0.20	35.3	-0.01	0.04	18.8
	clay fraction	161	-0.14	0.06	-0.16	0.08	-0.26	0.10	42.2	0.00	0.01	-
13 BMB	Burgmagerbein; paleokarst depression filled with Miocene sediments (mainly limestone weathering											
	residuum)	44.6	0.27	0.11	0.38	0.33	0.54	0.15	9.4	-0.19	0.05	8.22
	clay fraction	137	-0.27	0.04	-0.32	0.05	-0.49	0.09	21.4	-0.10	0.02	-
14 BMB	Burgmagerbein; paleokarst channels filled with Oligocene sediments (mainly limestone weathering											
	residuum)	180	0.26	0.14	0.28	0.27	0.42	0.22	32.7	0.03	0.04	24.3
15 HRD	Heroldingen; OSM?, clayish carbonate-rich silt to fine-											
	grained sand	59.9	0.08	0.07	0.16	0.14	0.17	0.17	58.0	-0.30	0.03	11.5
16 HRD	Heroldingen; OSM?, white fine-grained quartz sand	4.6	-0.05	0.04	-0.04	0.07	-0.10	0.06	0.82	-0.44	0.02	0.95
23 MHR	Möhren; freshwater oligocene limestone	66.4	0.58	0.12	0.80	0.31	1.10	0.19	12.2	0.19	0.02	7.97
24 RTS	Rothenstein; clay-rich paleokarst depression fill	118	-0.07	0.03	-0.12	0.29	-0.19	0.03	33.1	-0.35	0.04	30.9
25 RTS	Rothenstein; quartz-rich red depression fill	186	0.32	0.03	0.43	0.29	0.56	0.09	40.0	-0.17	0.02	25.2
26 LMH	Lohrmanshoff; dolomitized limestone	8.7	0.64	0.05	0.87	0.41	1.24	0.03	1.8	0.68	0.02	0.68
28 ALT	Altisheim; OSM, quartz-dominated sand with minor											/
	clay and carbonate	33.2	0.07	0.06	0.18	0.05	0.19	0.09	3.0	-0.15	0.01	5.26
Moldavites												
SBM-11	South Bohemia	14.9	2.23	0.20	3.32	0.40	4.46	0.37	0.42	5.35	0.07	10.9
SBM-23	South Bohemia	30.7	1.66	0.11	2.35	0.41	3.23	0.19	1.23	5.10	0.04	9.47
SBM-35	South Bohemia	25.9	3.20	0.11	4.69	0.37	6.28	0.20	0.63	4.82	0.05	8.50
SBM-44	South Bohemia	6.9	2.94	0.28	4.65	0.49	6.18	0.30	0.17	3.92	0.08	7.79
SBM-88	South Bohemia	34.5	1.82	0.12	2.64	0.31	3.55	0.17	1.44	5.02	0.10	12.2
SBM-192	South Bohemia	1.85	3.30	-	5.27	-	6.95	-	0.19	2.19	0.08	8.70
MM-60	Moravia	18.8	2.74	0.16	4.05	0.39	5.40	0.25	0.62	8.81	0.06	11.2
MM-67	Moravia	7.3	2.11	0.05	3.08	0.63	4.22	0.26	0.75	5.77	0.04	10.5
CHBM-5	Cheb Basin	108	1.96	0.15	2.86	0.28	3.82	0.23	2.86	1.62	0.04	8.70
CHBM-6	Cheb Basin	71.3	3.65	0.10	5.38	0.26	7.16	0.14	1.27	4.21	0.18	10.2

CHBM-7	Cheb Basin	58.7	3.32	0.19	5.03	0.27	6.58	0.31	0.39	12.5	0.01	9.62
MCB-2	Cheb Basin	20.6	2.32	0.23	3.56	0.26	4.68	0.35	1.83	2.72	0.05	-
LM-1	Lusatia	16.7	2.47	0.08	3.69	0.32	4.95	0.14	1.13	6.08	0.04	11.8
Reference materials												
BHVO-2			0.31	0.03	0.53	0.07	0.63	0.07		0.10	0.04	
BCR-2			0.17	0.05	0.28	0.11	0.34	0.10				

A double-pass quartz spray chamber combined with a 100 μ L/min nebulizer was used. Typical signals of ~3V for ⁶⁴Zn and ~10V for ⁶⁵Cu were achieved. For Zn, isobaric interference at mass 64 from Ni was found negligible by monitoring ⁶²Ni signal (<8×10⁻⁵ V) and assuming natural Ni isotope composition; levels of Ga and Cu were always lower than 2 mV and 20 mV, respectively.

A single measurement has only been performed for moldavite SBM-192 due to extremely low Zn content and a limited sample mass available for analysis.

^a Al₂O₃ contents from Žák et al. (2016), save for 26 LMH (this study; recalculated to volatile-free basis according to Žák et al., 2016).

Table 2: Statistical parameters for Zn isotope data								
	а	L	U	b	L	U		
	SMATR, m	ajor axis (M	A) regressio	n				
S	0.0029	-0.0296	0.0353	1.9077	1.8038	2.0214		
М	-0.1620	-0.5788	0.2548	2.0658	1.9184	2.2325		
M1	-0.0350	-0.3846	0.3147	2.0062	1.8788	2.1480		
M2	0.0450	-0.1533	0.2433	1.9629	1.8882	2.0424		
Т	0.0486	-0.1732	0.2705	2.0934	1.9651	2.2360		
	SMATR, M	A, zero inte	rcept (line fo	orced throug	h origin)			
S	0			1.9149	1.8468	1.9870		
М	0			2.0068	1.9727	2.0419		
M1	0			1.9932	1.9642	2.0228		
M2	0			1.9798	1.9622	1.9977		
Т	0			2.1208	2.0708	2.1728		
	Bootstrap, N	A with resa	ampled resid	uals				
S	0.0027	-0.0268	0.2840	1.9087	1.8171	2.0024		
М	-0.1665	-0.4993	0.1716	2.0674	1.9462	2.2029		
Т	0.0468	-0.1506	0.2445	2.0946	1.9753	2.2190		

Lines fitted to Zn isotope data. Intercepts (*a*), slopes (*b*) and their lower (L) and upper (U) 95% confidence limits, for sediments (S), moldavites (M) and other tektites (T). The groups M1 and M2 are cleaned M data (see text for explanation).

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Figure captions

Figure 1

Copper vs. Zn concentrations (a) and isotope compositions (b) in central European tektites and sediments from the Ries area. Clay fractions are also plotted. Copper shows significantly larger scatter in isotope compositions relative to Zn. The Cu and Zn elemental and isotope data for other tektites (gray field) and moldavites (dashed area) are from Moynier et al. (2009a, 2010).

Figure 2

The Zn contents versus Al_2O_3 contents in moldavites and sediments from the Ries area. A broadly positive correlation is observed for sediments whereas tektites show rather uniform Al_2O_3 contents at a large range of Zn abundances. Zinc and Al data for other tektites and moldavites are given in Moynier et al. (2009a).

Figure 3

The plot of δ^{68} Zn versus δ^{66} Zn for moldavites, sediments and clay fractions. The slopes for different groups (sediments and moldavites – this study, tektites – Moynier et al., 2009a) imply similar processes of Zn isotope fractionation for samples modified at high temperatures (kinetic effects) and for those that underwent only low-temperature consolidation (equilibrium effects); see main text for further discussion. Dashed circle denotes australite 5772 core (Moynier et al., 2009a) which is clearly off the linear trend and also shows peculiar Zn enrichment (265 ppm). Samples SBM-44 and SBM-192 are marked and were discussed in terms of statistical evaluation (see Section 4.1).

Figure 4

The plot of Zn (a) and Cu (b) abundances and isotope compositions versus possible paths of a Rayleigh distillation process. Several starting points for Zn and Cu abundance were used to illustrate the concentration range of these elements in sedimentary materials of the Ries area.

Rayleigh distillation paths for Cu(I) species are nearly identical to Cu(II) species and a case for Cu_2O is only plotted. The primitive upper mantle and the upper continental crust values are from McDonough and Sun (1995) and Rudnick and Gao (2014).







