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2	North-American microtektites are more oxidized than tektites.							
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25	Abstract							
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27 Iron oxidation states and coordination numbers have been determined by micro-28 X-ray Absorption Near Edge Spectroscopy (XANES) on the cores of a large group of 29 microtektites from the Australasian, Ivory Coast, and North American (NA) tektite 30 strewn fields. The North American microtektites used in this study have been 31 collected from five sites at different distances from the source crater; most have SiO_2 content between 70 and 80 wt%. Accurate analysis of the pre-edge peak energy 32 position and integrated area allowed determination of $Fe^{3+}/(Fe^{2+}+Fe^{3+})$ ratios on all 33 34 samples with an estimated error of ± 0.05 .

Microtektites from the Australasian and Ivory Coast strewn fields show low values of the $Fe^{3+}/(Fe^{2+}+Fe^{3+})$ ratios, in fair agreement with tektites from the same strewn field. In contrast, microtektites from the North American strewn fields show a wide range of $Fe^{3+}/(Fe^{2+}+Fe^{3+})$ ratios from 0.02 to ca. 0.61. Comparison of Fe oxidation state data with chemical composition do not show any relation between $Fe^{3+}/(Fe^{2+}+Fe^{3+})$ ratios and Na, Ca, or K contents, thus suggesting that the high Fe oxidation states are not the consequence of sea-water alteration.

The difference between the Fe oxidation state of tektites and microtektites from the North American strewn fields suggests that some factors in the formation of the North American microtektites were different than for the North American tektites and for microtektites in the other strewn fields.

Previous Fe oxidation state data on NA tektites strongly suggest that the wide range in Fe oxidation state we found on NA microtektites is not related to lateral heterogeneity of the target rocks. Despite a correlation between microtektite oxidation state and distance from the source crater, we maintain that Fe oxidation state is not related only to the microtektite droplet flight distance. This is in keeping with the fact that no significant variations in the Fe oxidation state have been found in

52	microtektites from the Australasian strewn field, even for Australasian microtektites
53	recovered in Antarctica. The Fe oxidation state in North American microtektites could
54	be explained by interaction of melt droplets with a H ₂ O-rich vapor plumes generated
55	during the impact. These data point out that some difference must exist between the
56	thermal histories of microtektites and tektites from the NA strewn field. Moreover,
57	microtektites from the NA strewn field show also distinctively higher oxidation states
58	than those from Ivory Coast or the Australasian strewn fields.
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60	Keywords: impact glasses, tektites, microtektites, Fe local structure, XANES
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62	INTRODUCTION
63	Microtektites are small (<1 mm) splash-form impact glasses related to tektites,
64	which are found scattered over regions of the Earth's surface called strewn fields (e.g.
65	Koeberl, 1986, 1994). They are associated with three of the four tektite strewn fields
66	so far known: the North American (NA), the Ivory Coast, and the Australasian strewn
67	fields. They are generally found in deep sea cores (e.g., Glass, 1967, 1968, 1972;

Cassidy et al., 1969), and their distribution greatly contributed to establish the limits
of tektites strewn fields (see Glass, 1990; Glass and Pizzuto, 1994).

Despite the availability of geochemical studies on microtektites (see Glass et al, 2004 and references therein), few studies exist of the Fe coordination number and oxidation state in such materials (Senftle et al., 1969). As microtektites constitute a large fraction of the mass of the glass produced by a tektite-generating impact event, such studies are of great importance for a more complete understanding of impactgenerated glasses and, in particular, to try reconstructing the oxygen fugacity conditions prevailing during impact melt formation. 77 Previous data have shown a set of microtektites from the North American strewn field (collected at the DSDP Site 94) to be consistently more oxidized with respect to 78 79 microtektites from the other strewn fields (Giuli et al., 2008b). Moreover, another 80 study of macroscopic tektites from the North-American strewn fields showed no 81 variation of the Fe oxidation state as a function of textite provenance and composition 82 (Giuli et al., 2010a), thus confirming that the strong variation in the Fe oxidation state 83 previously found in NA microtektites does not depend on lateral heterogeneities of 84 the target rock.

85 This case is unique among tektites and microtektites and, if confirmed, may 86 contribute to a better understanding of microtektite formation processes. To expand 87 on our previous data, we studied a larger set of microtektites from the North 88 American strewn field collected from three additional cores (RC9-58 and DSDP sites 89 612 and 149) and from Barbados. These samples have been collected at different 90 distances from the source crater and span a wider compositional range than those 91 previously analyzed. The use of a small X-ray beam allowed to probe the core of the 92 studied samples, thus avoiding possible alteration rims.

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SAMPLES AND EXPERIMENTAL

Microtektite samples consist of glassy spherules retrieved from deep-sea cores from three different strewn fields: Australasian microtektites (including 1 normal, 3 high Ni, 1 high Mg, and 1 intermediate sample) have been sampled from ODP Site 769A, and cores RC12-331 and RC14-46; Ivory Coast microtektites (10 samples) come from core K9-56; North American microtektites (25 samples) come from DSDP Sites 612, 94, and 149, core RC9-58, and from Bath Cliff, Barbados. The compositions of these samples have already been studied (Glass et al., 2004) and are

reported in Table 1. The spherules have been embedded in resin, ground down to expose an interior surface, and polished. Typical dimensions range from 400 to 800 μm diameter. Both scanning electron microscopy and optical microscopy indicate that no crystals are present. As shown in Fig. 1, most of the microtektite samples display silica contents ranging between ca. 70 and 80 wt % and alkali contents ranging between ca. 3 and 5 wt %.

108 The standards used for X-ray absorption near edge spectroscopy (XANES) 109 measurements are: a staurolite from Canton Ticino (Switzerland) and a synthetic Feakermanite for Fe^{2+} in tetrahedral coordination: a grandidierite from Madagascar for 110 Fe^{2+} in trigonal dipyramidal coordination; a synthetic kirschsteinite, a synthetic 111 orthopyroxene ($En_{60}Fs_{40}$) and a siderite from Erzberg (Austria) for Fe^{2+} in octahedral 112 coordination; an andradite from Italy, and an aegirine from Malawi for Fe^{3+} in 113 octahedral coordination; a voderite from Mautia Hills (Tanzania) for Fe³⁺ in 5-fold 114 coordination; a tetra-ferriphlogopite from Tapira (Brasil) for Fe^{3+} in 4-fold 115 116 coordination (Giuli et al., 2001). The natural standards were separated by hand-117 picking from cm-sized crystals choosing the clearest portions to avoid impurities. All 118 the standards were checked for purity by both optical microscopy and X-ray 119 diffraction. Model compounds for X-ray Absorption Spectroscopy (XAS) 120 measurement were prepared by smearing finely ground powder on a kapton tape, 121 whereas the glass samples were prepared as slides with polished surfaces. Fe K-edge 122 XANES spectra were recorded at room temperature at the ESRF (European 123 Synchrotron Radiation Facility) storage ring on the undulator beamline ID26 124 (Gauthier et al., 1999; Solé et al., 1999) operating at 6 GeV. A Si(311) double-crystal 125 monochromator was used, providing an energy resolution of ~ 0.2 eV at the Fe K-126 edge. However, the main limitation for energy resolution is the finite core-hole width

127	of the absorbing element (~1.15 eV at the Fe K-edge, Krause and Oliver 1979),
128	resulting in a convoluted energy resolution of \sim 1.4 eV. The energy was calibrated by
129	defining the first derivative peak of a metallic Fe reference foil to be at 7112.0 eV.
130	Two Si mirrors were used for the harmonics rejection of the incident X-ray beam.
131	Beam dimension at the sample was 55x120 micrometers. For each sample the beam
132	has been carefully centered by means of vertical and horizontal scans so as to allow
133	probing the inner part of the spherules far from the border, thus avoiding possible
134	alteration rims. XANES data were recorded in quick-scan mode by simultaneously
135	scanning the monochromator angle and the undulator gap with a typical energy step
136	of 0.1 eV and counting 44 ms per point. Each scan took 120 seconds and an average
137	of 12 spectra was taken per sample. The spectra were acquired in fluorescence mode,
138	using a Si photo-diode, and I_0 was monitored by measuring the fluorescence signal of
139	a titanium foil using a Si photo-diode. The flat surface of the sample was positioned at
140	45° with respect to the beam and the detector. Average of about 12 scans for each
141	sample allowed us to obtain very good signal to noise ratios despite the small beam
142	size. The energy reproducibility has been estimated to be ± 0.03 eV or better.
143	Experimental XANES spectra were reduced by background subtraction with a

143 144 linear function and then normalized for atomic absorption on the average absorption 145 coefficient of the spectral region from 7170 to 7350 eV. The threshold energy was 146 taken as the first maximum of the first derivative of the spectra, whereas peak 147 positions were obtained by calculating the second derivative of the spectra. Pre-edge 148 peak analysis was carried out following the same procedure reported in Wilke et al. 149 (2001) and Giuli et al. (2002). The pre-edge peak was fitted by a sum of pseudo-Voigt 150 functions, and their intensities along with energy positions were compared with those 151 of the standards analysed here and others from the literature (e.g., Wilke et al., 2001;

152 Farges, 2001) in order to extract information on Fe oxidation state and coordination 153 number in these microtektites. Particular care was taken in using the smallest possible 154 number of components in the pre-edge peak fitting procedure. In particular, the 155 number (and, in some cases, the approximate energy) of the components was 156 constrained to equal the number (and approximate energy) of the minima in the 157 second derivative spectrum of the pre-edge peak. Several different procedures have 158 been attempted for peak fitting: simultaneous fitting of the background and the 159 pseudo-Voigt component; preliminary background subtraction followed by fitting 160 with pseudo-Voigt component allowing the components to have different full width at 161 half maximum (FWHM) or Lorentian character; preliminary background subtraction 162 followed by fitting with pseudo-Voigt component constrained to have the same 163 FWHM and Lorentian character. This last procedure has been shown to produce the 164 best results in terms of data scatter.

165 The precision of the fitted pre-edge peak centroid energy and integrated area are 166 ± 0.02 eV and ± 0.015 , respectively. Mixing lines have been calculated as linear combination of two pre-edge peaks representative of Fe^{2+} and Fe^{3+} (see Giuli et al., 167 2003, 2011). We tried different integrated areas for Fe^{2+} and Fe^{3+} until a calculated 168 169 mixing line was found to match the integrated area and centroid energy of a set of glasses; then it has been used to evaluate the contribution of the divalent and trivalent 170 171 Fe to the experimentally measured pre-edge peaks. This procedure implies that a 172 single spectrum (pre-edge peak integrated area and centroid energy) does not allow an accurate determination of the $Fe^{3+}/(Fe^{2+}+Fe^{3+})$ ratio unless contrains are applied on 173 the integrated areas relative to the Fe^{3+} and Fe^{2+} species contributing to that spectrum. 174 175 This procedure, applied to a set of synthetic glasses (Giuli et al., 2011), resulted in accurate determinations of $Fe^{3+}/(Fe^{2+}+Fe^{3+})$ ratios that are in agreement with 176

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177 independent titration data within \pm 0.03. In the present case we estimate the error in 178 the determination of Fe³⁺/(Fe²⁺+Fe³⁺) ratios on microtektite samples to be \pm 0.05

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XANES RESULTS

181 Examples of experimental Fe K-edge XANES spectra are shown in Fig. 2 for five 182 groups of microtektites: Australasian and Ivory Coast in Fig. 2a and 2b, respectively, 183 and three groups of North American microtektites in Fig. 2 c, d, and e. The shape and 184 edge energy position of the Australasian and Ivory Coast microtektite spectra are very 185 similar to those of splash form tektites studied previously (Giuli et al., 2002), 186 suggesting the presence of predominantly divalent Fe. On the other hand, the spectra 187 of the North American microtektite samples display significant differences in both the 188 shape of the XANES spectra and their first derivative that are clearly related to 189 variations in the Fe oxidation state. Moreover, also significant changes in the pre-edge 190 peak (labeled P) are visible.

191 The pre-edge peak is related to an s-d like electronic transition and, despite being 192 dipole-forbidden, can become partially allowed by mixing of the d-states of the 193 transition metal with the *p*-states of the surrounding oxygen atoms. This means that 194 the pre-edge peak energy position and intensity depend strongly on both the geometry 195 around Fe and on the mean Fe oxidation state (Calas and Petiau, 1983; Brown et al., 196 1995). As already shown in the literature, accurate evaluation of the pre-edge peak 197 centroid energy and integrated area and comparison with those of Fe model 198 compounds can provide quantitative information on both Fe oxidation state and 199 coordination environment (see Calas and Petiau, 1983; Brown et al., 1995; Wilke et 200 al., 2001, 2006; Farges, 2001; Giuli et al., 2002, 2011): its intensity will be almost

201 zero in case of regular octahedral symmetry (O_h) around the absorber, whereas it will

202 reach its maximum in the case of tetrahedral simmetry (T_d).

203 The background subtracted pre-edge peaks of the microtektite XANES spectra are 204 shown in Fig. 3 (a-e), along with the pseudo-Voigt components used in the fitting 205 procedure and their sums, whereas their centroid energies and integrated areas are 206 reported in Table 2. Each pre-edge peak can be fitted with two to three components 207 whose energies (ca. 7112.6, 7113.9-7114.3, and 7114.5-7115.3 eV) are consistent 208 with those of divalent and trivalent Fe model compounds. In particular, while the first and last components can be ascribed to contributions from Fe^{2+} and Fe^{3+} respectively. 209 the component at intermediate energy results from contributions by both Fe^{2+} and 210 Fe³⁺. The relative importance of divalent or trivalent Fe causes an increase of the 211 212 respective components resulting in an energy shift of the pre-edge peak centroid.

213 The integrated area of the pre-edge peaks vs. their centroid energies are plotted in 214 Figure 4 (a,b) along with the data of Fe model compounds analysed here and by 215 others (Wilke et al., 2001; Farges, 2001; Giuli et al., 2002). For the sake of simplicity, 216 the relative energy is plotted (0 refers to the first maximum of the first derivative of 217 metallic Fe spectrum) in order to avoid confusion when comparing data with literature 218 data where a different energy value of metallic Fe has been chosen. All divalent Fe 219 model compounds plot at energies close to 0.9 eV above the metallic Fe edge, 220 whereas trivalent Fe model compounds plot at energies close to 2.4 eV. At constant 221 energy, the intensity of the model compounds pre-edge peaks varies according to the 222 Fe coordination geometry (the shaded ellipses refer to the range of coordination 223 numbers in Fe model compounds).

The Australasian and Ivory Coast microtektites data (Fig. 4a) plot within a narrow region close in energy to that of divalent Fe model compounds, meaning that most of

the Fe is divalent. Three mixing lines (dotted line with small diamonds) have been

- 227 calculated by a linear combination of:
- i) a pre-edge peak with a centroid at 0.9 eV above the edge of metallic iron and an integrated area intermediate to that of ${}^{[4]}Fe^{2+}$ and ${}^{[5]}Fe^{2+}$ model compounds;
- ii) a pre-edge peak with a centroid at 2.4 eV above the edge of metallic iron and
- integrated area typical of ${}^{[4]}Fe^{3+}$, or ${}^{[5]}Fe^{3+}$, or ${}^{[6]}Fe^{3+}$.

These mixing lines stem from a region with intermediate intensity between $^{[4]}Fe^{2+}$ and $^{[5]}Fe^{2+}$ thus meaning that the divalent Fe in these glasses is present both in [4] and [5] coordination. Should $^{[6]}Fe^{2+}$ be present in consistent amounts it would drastically lower the pre-edge peak intensity. However, its presence in minor amounts cannot be ruled out by XANES data alone. The two mixing lines that best fit the experimental points are those calculated using $^{[4]}Fe^{3+}$ and $^{[5]}Fe^{3+}$ pre-edge peak intensity; this suggests that trivalent iron may be in [4] and [5] coordination.

Comparison between the experimental pre-edge peak data and the calculated mixing lines provided quantitative $Fe^{3+}/(Fe^{3+}+Fe^{2+})$ ratios $\leq 0.1 \pm 0.05$ for most of the samples and up to 0.21 for four samples (0.13, 0.15, 0.19, and 0.21). Based on the precision of the energy (± 0.03 and ± 0.05 eV for the Ivory Coast and Australasian spectra respectively), we estimate the error in the $Fe^{3+}/(Fe^{3+}+Fe^{2+})$ ratios to be within ± 0.05 . These results are in keeping with known Fe oxidation states of Australasian and Ivory Coast microtektites.

On the other hand, North American microtektites display centroid energy values much higher than those of the Australasian and Ivory Coast microtektites, indicating significantly higher contributions from trivalent Fe (Fig. 4b). Three mixing lines have been calculated as in the previous case. It is worth remarking that this set of mixing lines originates from a point with the average intensity of a wide set of tektites

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251 analysed so far (Giuli et al., 2002, 2010a, 2010b). As in the Ivory Coast and 252 Australasian microtektite case, the average coordination number of divalent Fe is 253 intermediate between [4] and [5]. However, while many the samples can be explained as consisting of a mixture of ${}^{[4]}Fe^{2+}$, ${}^{[5]}Fe^{2+}$, ${}^{[4]}Fe^{3+}$, ${}^{[5]}Fe^{3+}$, (see Calas et al., 1983; 254 255 Brown et al., 1995; Rossano et al., 2009; Wilke et al., 2004, 2006; Jackson et al., 256 2005; Giuli et al., 2002, 2011; Weigel et al., 2006, 2008), which are common Fe 257 species usually found in silicate melts and glasses, the microtektites from Barbados have distinctly lower integrated areas, compatible with the presence of ${}^{[6]}Fe^{3+}$ along 258 259 with other Fe species. Despite this Fe species not being very common in silicate melts 260 and glasses, it has been observed by Wilke et al. (2004) in synthetic glasses of basaltic 261 composition. Comparison between the experimental pre-edge peak data of NA microtektites and the calculated mixing lines provided quantitative $Fe^{3+}/(Fe^{3+}+Fe^{2+})$ 262 263 ratios ranging from 0.02 and 0.61 ± 0.05 .

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DISCUSSION

266 XANES data have been presented for a large set of microtektites from the 267 Australasian, Ivory Coast and North American strewn fields (6, 10, and 26 samples 268 respectively). Accurate analysis of the pre-edge peak allowed determination of the Fe 269 oxidation state for all the samples analyzed (see Table 2). To the authors' knowledge 270 this is the first report of the Fe oxidation state in such a large set of microtektites.

271 Most of the samples from the Australasian and Ivory Coast strewn fields display values consistent with $Fe^{3+}/(Fe^{2+}+Fe^{3+})$ ratios $\leq 0.1 \pm 0.05$, with only four samples 272 273 displaying slightly larger values (0.13, 0.15, 0.19, and 0.21). Thus, in these two 274 strewn fields, microtektites display Fe oxidation states and coordination numbers 275 fairly similar to those of macroscopic tektites (Fudali et al., 1987; Dunlap et al, 1998;

276 Rossano et al., 1999; Giuli et al., 2002, 2010a, 2010b). On the other hand, 277 microtektites from the North American strewn field exhibit much higher values of the 278 $Fe^{3+}/(Fe^{2+}+Fe^{3+})$ ratios, from 0.02 up to 0.61 ± 0.05.

279 As there is the possibility that a long residence time in marine sediments may have 280 altered the microtektite glasses, we checked whether some correlation may exist 281 between Fe oxidation state and chemical composition which may be indicative of sea-282 water alteration induced oxidation (see Giorgetti et al., 2001; Alt and Mata, 2000). In Fig. 5 the alkali content is plotted as a function of the $Fe^{3+}/(Fe^{2+}+Fe^{3+})$ ratio of all the 283 284 microtektites analyzed here. No obvious correlation has been found between Fe 285 oxidation state and chemical composition of the studied microtektites. In particular, 286 no inverse relationship has been found with the Na content, which might have been 287 expected in case of oxidation due to sea-water alteration, nor any other correlation has 288 been found that would suggest alteration in a marine environment. Moreover, a tektite 289 fragment retrieved along with microtektites from a core in the DSDP Site 612 (and which, thus, should have similar degree of alteration) displayed an $Fe^{3+}/(Fe^{2+}+Fe^{3+})$ 290 291 ratio of 0.05 ± 0.05 (Giuli et al., 2010a), consistent with values for the textites found 292 on land, thus further suggesting that there was no alteration-induced oxidation in these 293 microtektite samples. Also preliminary micro-infrared spectroscopy data on a few 294 microtektite samples (see supplementary material) yielded low water contents in the 295 range between 35 to 455 ppm, similar to those of NA tektites (see Beran and Koeberl, 296 1997; Giuli et al., 2010c (IMA abstract)), reinforcing the suggestion that the 297 microtektites analyzed here did not suffer alteration.

Thus, we believe that the oxidation states determined are a signature of the formation process of these NA microtektites, and not the product of subsequent alteration in the sediment. Interestingly, pre-edge data of all NA microtektites seem to define different trends all originating from a single narrow region whose energy and intensity is typical of most splash form tektites studied so far. Also, previous data on NA tektites (Giuli et al., 2010a) from different locations and with different chemistry clearly show that the Fe oxidation state of tektites does not vary according to tektite composition. This indicates that the wide range of Fe oxidation states we found in NA microtektites is not related to lateral heterogeneity of the target rock.

A possible explanation may involve formation of microtektites with Fe oxidation state and coordination number similar to that of tektites and Trinity glass (a glass formed by high temperature melting of desert sand during the first atomic bomb test at the Trinity site; see Glass et al., 1988; Giuli et al., 2010b), and subsequent oxidation when still molten.

Although a similarity with impact glass spherules from the K/T boundary, which showed a similar trend in pre-edge peak data extending up to 100% trivalent Fe, may be envisioned (Giuli et al., 2005, 2008a), no clear explanation can be provided yet for the high Fe oxidation state of the North American microtektites.

316 In this respect it is interesting to notice that the Fe oxidation state of the studied 317 microtektites seems to increase with the distance of the collection site from the source 318 crater. In Fig. 6 the fraction of the trivalent Fe is shown as a function of the distance 319 of the microtektite collection site from the Chesapeake Bay impact structure: even 320 though microtektites within every collection site display a range of Fe oxidation 321 states, it is evident that the fraction of trivalent Fe tends to increase with the distance 322 from the crater. Although it would be tempting to relate the increase of the trivalent 323 Fe fraction to the oxidation of melt in air during the travel from the source crater to 324 the landing site, this hypothesis has to be discounted when comparing data for 325 Australasian microtektites with those found in Antarctica (Folco et al., 2010). In fact,

326 recent analyses of the Fe oxidation state on 29 Antarctic microtektite samples (Giuli 327 et al., 2012) showed that the fraction of trivalent Fe is lower than 0.1, thus being 328 compatible with the Fe oxidation state of Australasian tektites previously analyzed 329 (see Dunlap et al. 1998; Dunlap and Sibley, 2004; Rossano et al., 1999; Giuli et al., 330 2002) and the microtektites analyzed here. In that case, a flight distance of up to ca. 331 11000 km from the putative location of the source crater did not produce any 332 noticeable oxidation of the Fe in the microtektite melt droplet; thus, oxidation in air 333 during the flight is unlikely to be responsible for the higher Fe oxidation state of NA 334 microtektites compared to tektites.

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IMPLICATIONS

337 Even though the Fe oxidation in North American microtektites could be explained 338 by interaction of melt droplets with a H₂O-rich vapor plumes generated during the 339 impact, no definitive explanation can yet be given for the anomalously high Fe 340 oxidation state of NA microtektites and more studies are needed to understand the 341 factors affecting the Fe oxidation state of these samples. However, the marked difference in the $Fe^{3+}/(Fe^{2+}+Fe^{3+})$ ratio of tektites and microtektites from the North 342 343 American strewn field raises the question whether or not microtektites should simply 344 be considered as microscopic analogous of tektites or if some difference exists 345 between their formation mechanisms. Furthermore, our results indicate that some 346 differences must exist in the thermal (and/or oxygen fugacity) histories of 347 microtektites from the NA strewn field compared with those from the Ivory Coast or 348 the Australasian strewn fields.

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503 **Figure captions**

504

505	Fig. 1 Total alkalis versus silica contents of the analyzed North American
506	microtektites. For source of compositional data see Table 1. The open rectangle
507	represents the range in the silica and alkali contents for North American tektites as
508	reported by Glass et al. (1995, 1998) and Montanari and Koeberl (2000). Each sample
509	group was given the designation NA plus the sample name (3 digit number) as
510	reported in Glass et al. (2004) and the collection site.

511

Fig. 2: Experimental Fe K-edge XANES spectra of microtektites: a) Ivory Coast; b) Australasian; c) North American (DSDP 612); d) North American (DSDP 149); e) North American (Barbados). Pre-edge peak is labeled "P". The spectra labels are the same as sample names in Table 1. The first three digits refers to the disc number the microtektites were mounted on and the number after the hyphen refers to the number of the microtektite on that disc.

518

519 Fig. 3: Background subtracted pre-edge peaks: a) Ivory Coast; b) Australasian;
520 c) North American (DSDP Site 612); d) North American (DSDP Site 149); e) North
521 American (Barbados). Labels as in Fig. 2.

522

Fig. 4: Pre-edge peak integrated area versus centroid energy of : a) Ivory Coast
and Australasian microtektites; b) North American microtektites. Symbols as in Fig.
1. The 0 on the x axis refers to the edge energy of metallic iron (7112.0 eV).

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- 527 Fig. 5: Alkali content (oxide wt %) as a function of the fraction of trivalent Fe
- 528 in the studied microtektites. The amount of both Na₂O and K₂O does not show any
- 529 significant decrease as would be expected in case of sea water alteration.
- 530
- 531 Fig. 6: Trivalent Fe fraction of the NA microtektites as a function of collection
- 532 site distance from the source crater (symbols as in Fig 4b). A marked increase of the
- 533 trivalent Fe at increasing distance from the crater is evident.
- 534
- 535

536 Fig.1



539 Fig. 2





543 Fig. 3





547 Fig. 4a



548





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555 Fig.6



Sample	core	SiO ₂	Al ₂ O ₃	Cr ₂ O ₃	FeO	CaO	MgO	K ₂ O	Na ₂ O	TiO ₂	total	Source [†]
Australasian microtektites												
805-1	ODP769A	64.72	16.21	0.03	5.39	3.9	7.77	0.48	0.62	0.87	99.99	1
805-2	ODP769A	70.09	13.54	0.01	5.74	2.68	3.02	2.77	1.37	0.75	99.97	1
805-3	ODP769A	66.84	16.71	0.01	5.65	2.72	2.62	3.21	1.39	0.83	99.98	1
805-4	RC12-331	56.05	20.66	0.05	4.5	5.72	11.34	0.13	0.43	1.17	100.05	1
805-5	RC14-46	67.17	14.94	0.01	5.97	3.68	3.07	2.79	1.59	0.8	100.02	1
805-6	RC14-46	69.25	12.48	0.04	6.77	3.29	4.17	2.22	1.04	0.69	99.95	1
Ivory Coast microtektites												
858-1	K9-56	63.68	17.62	0.05	7.44	2.09	6.15	1.03	1.67	0.57	100.30	1
858-2	K9-56	69.30	15.82	0.02	5.67	1.48	3.25	1.89	2.10	0.47	100.00	1
858-3	K9-56	67.35	17.15	0.01	6.39	1.56	3.13	1.90	1.98	0.54	100.01	1
858-4	K9-56	67.33	16.50	0.02	6.48	1.63	3.86	1.65	1.96	0.58	100.01	1
858-5	K9-56	67.12	15.97	0.02	6.53	1.88	4.20	1.82	1.96	0.50	100.00	1
858-6	K9-56	67.38	16.89	0.02	6.31	1.54	3.31	1.96	2.09	0.52	100.02	1
858-7	K9-56	68.11	17.02	0.02	6.67	1.05	2.91	2.03	1.60	0.54	99.95	1
858-8	K9-56	67.86	16.23	0.02	6.34	1.81	3.84	1.54	1.80	0.53	99.97	1
858-9	K9-56	68.00	16.90	0.02	6.55	1.14	3.15	1.95	1.72	0.58	100.01	1
858-10	K9-56	67.59	16.73	0.02	6.19	1.54	3.37	1.96	2.03	0.54	99.97	1
				No	rth Ame	rican mic	rotektites					
772-1	DSDP94	71.44	17.62	0.05	5.98	0.43	0.95	1.64	1.05	0.90	100.06	1
772-2	DSDP94	77.87	11.66	0.02	3.09	1.29	1.10	3.16	1.22	0.60	100.01	1
772-3	DSDP94	74.36	13.82	0.03	3.51	2.01	1.68	2.60	1.25	0.74	100.00	1
772-4	DSDP94	78.47	11.78	0.02	3.12	1.30	1.23	2.52	0.94	0.60	99.98	1
772-5	DSDP94	73.60	13.87	0.05	4.17	1.78	1.76	2.74	1.32	0.76	100.05	1
772-6	DSDP94	81.51	10.82	0.01	2.28	0.45	0.80	2.74	0.92	0.46	99.99	1
772-7	DSDP94	79.97	11.03	0.02	2.78	0.94	0.96	3.02	0.78	0.51	100.01	1
392-1	DSDP612	71.01	16.10	n.d.	4.16	1.61	1.23	1.40	3.01	0.85	99.37	3
392-2	DSDP612	67.38	15.50	n.d.	5.43	2.63	3.49	1.52	2.65	0.75	99.35	2
392-6	DSDP612	69.94	15.10	n.d.	4.33	2.11	2.60	1.51	3.19	0.65	99.43	2
151-91a	RC9-58	75.19	12.73	n.d.	3.04	1.52	1.30	4.07	1.28	0.46	99.59	3
151-93b	RC9-58	67.52	16.06	n.d.	5.51	1.42	1.69	4.84	1.45	1.07	99.56	3

Table 1. Composition of the studied microtektites in wt% oxides.

151-94a	RC9-58	79.63	11.31	n.d.	2.59	0.52	1.00	2.95	1.02	0.53	99.55	3
151-102a	RC9-58	76.12	13.37	n.d.	3.62	0.97	1.00	2.99	0.92	0.66	99.65	3
775-1	DSDP149	71.38	14.57	n.d.	4.29	1.55	1.95	4.30	1.24	0.73	100.01	1
775-2	DSDP149	72.22	15.63	n.d.	4.90	1.15	1.48	2.86	0.89	0.85	99.98	1
775-3	DSDP149	70.72	16.30	n.d.	4.68	1.56	1.68	3.20	1.01	0.84	99.99	1
312-2	Barbados	75.40	13.20	n.d.	4.95	0.76	1.00	2.31	0.92	0.60	99.14	4
312-3	Barbados	75.40	13.60	n.d.	3.58	1.26	0.97	2.70	1.02	0.65	99.18	4
312-4	Barbados	78.90	13.70	n.d.	1.56	1.88	0.70	1.55	0.21	0.61	99.11	4
312-5	Barbados	77.80	12.40	n.d.	2.78	1.03	0.64	3.08	0.98	0.52	99.23	4
312-6	Barbados	76.10	13.30	n.d.	3.75	0.77	0.69	2.33	1.59	0.69	99.22	4
312-7	Barbados	73.80	14.20	n.d.	4.29	1.43	1.17	2.65	1.32	0.68	99.54	4

[†]1 = Glass et al. (2004); 2 = Glass (1989); 3 = this study; 4 = Burns (1985).

Sample name	Centroid (eV) ^a	Integrated intensity	Fit agreement index (%)	Fe ³⁺ /(Fe ²⁺ +Fe ³⁺)
Ivory Coast microtektites				
858-1	7112.98	0.149	99.97	0.03
858-2	7113.15	0.142	99.95	0.17
858-3	7113.08	0.141	99.96	0.12
858-4	7113.15	0.146	99.97	0.15
858-5	7113.21	0.157	99.97	0.12
858-6	7113.13	0.160	99.97	0.08
858-7	7113.16	0.161	99.97	0.09
858-8	7113.24	0.163	99.96	0.13
858-9	7113.22	0.155	99.97	0.15
858-10	7113.08	0.152	99.97	0.06
Australasian microtektites				
805-1	7112.93	0.141	99.92	0.02
805-2	7113.28	0.143	99.98	0.22
805-3	7112.98	0.137	99.96	0.04
805-4	7113.13	0.139	99.95	0.13
805-5	7113.23	0 144	99.92	0.19
805-6	7113.03	0.143	99.92	0.07
North American microtektites	/115.05	0.115	<i>)).)</i> 2	0.07
392-1	7112.94	0.130	99 98	0.02
392-2	7113.18	0.146	99 94	0.16
392-6	7113.10	0.149	99.95	0.12
775-1	7113.12	0.149	99.87	0.12
775-2	7113.55	0.136	99.94	0.24
775-3	7113.19	0.150	99.94	0.24
312-2	7113.40	0.128	99.87	0.50
312-3	7113.50	0.120	99.90	0.50
312-4	7113.01	0.123	99.89	0.54
312-5	7113.55	0.124	99.91	0.50
312-5	7113.71	0.131	99.92	0.58
312-0	7113.72	0.123	00 02	0.50
772 1	7113.30	0.127	00.06	0.32
772-1	7113.24	0.138	99.90	0.25
772.2	7113.30	0.138	99.00	0.33
772-3	7113.13	0.170	99.94	0.00
772-4	7113.32	0.143	99.07	0.42
772.6	7113.12	0.101	99.92	0.00
772-0	/113.14	0.131	99.94	0.22
1/2-7	/113.5/	0.133	99.85	0.49
151-91a 151-02-	/113.48	0.152	99.93	0.35
151-938	/113.54	0.161	99.95	0.36
151-930	/113.29	0.135	99.95	0.30
151-94a	7113.45	0.130	99.93	0.43
151-101b	7113.55	0.135	99.92	0.47
151-102a	7113.37	0.148	99.96	0.29

Table 2. Pre-edge peak features in the samples studied.

^a The energy has been calibrated by setting the Edge energy of metallic Fe at 7112.0 eV