Meteoroid atmospheric entry investigated with plasma flow experiments: Petrography and geochemistry of the recovered material

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A B S T R A C T

Melting experiments attempting to reproduce some of the processes affecting asteroidal and cometary material during atmospheric entry have been performed in a high enthalpy facility. For the first time with the proposed experimental setup, the resulting material has been recovered, studied, and compared with natural analogues, focusing on the thermal and redox reactions triggered by interaction between the melt and the atmospheric gases under high temperature and low pressure conditions. Experimental conditions were tested across a range of parameters, such as heat flux, experiment duration, and pressure, using two types of sample holders materials, namely cork and graphite. A basalt served as asteroidal analog and to calibrate the experiments, before melting a H5 ordinary chondrite meteorite. The quenched melt recovered after the experiments has been analyzed by μ-XRF, EDS-SEM, EMPA, LA-ICP-MS, and XANES spectroscopy.

The glass formed from the basalt is fairly homogeneous, depleted in highly volatile elements (e.g., Na, K), relatively enriched in moderately siderophile elements (e.g., Co, Ni), and has reached an equilibrium redox state with a lower Fe3+/Fe2+ ratio than that in the starting material. Spherical objects, enriched in SiO2, Na2O and K2O concentrations, were observed, inferring condensation from the vaporized material. Despite instantaneous quenching, the melt formed from the ordinary chondrite shows extensive crystallization of mostly olivine and magnetite, the latter indicative of oxygen fugacity compatible with presence of both Fe3+ and Fe2+. Similar features have been observed in natural meteorite fusion crusts and in micrometeorites, implying that, at least in terms of maximum temperature reached and chemical reactions, the experiments have successfully reproduced the conditions likely encountered by extraterrestrial material following atmospheric entry.

1. Introduction

Atmospheric entry of asteroidal and cometary material likely implies melting, evaporation, loss of mass, physical separation of metal-rich phases, and redox variations. All these processes may affect microstructures of and induce chemical changes in the recovered meteorites and micrometeorites. Ablation is an important factor, but if the meteoroid is large enough a portion survives and falls on the Earth.
covered by a quenched melt film called fusion crust (e.g., Genge and Grady, 1999a). Similar processes might also affect microscopic dust particles (10–2000 μm) that enter the Earth’s upper atmosphere and are recovered on Earth as micrometeorites (e.g., Genge et al., 2008; Rubin and Grossman, 2010), as locally similar features are observed, for example in the form of a melt film coating the surface (e.g., Toppani and Libourel, 2003; Genge, 2006). The degree of melting and therefore the preservation potential of internal features mainly depend on the physico-chemical parameters of the extraterrestrial material (i.e., size, density, composition) and on the entry conditions (i.e., velocity, angle; e.g., Love and Brownlee, 1991). Based on the degree of melting experienced during atmospheric passage by micrometeoroids, the distinction is generally made between fully melted micrometeorites (or cosmic spherules), partially melted micrometeorites (or scoriaeous MMs), and unmelted or angular micrometeorites (e.g., Maurette et al., 1994; Taylor et al., 1998; Genge et al., 2008). Thus, to some degree, analogies may exist between the processes that lead to the formation of cosmic spherules and those that generate meteorite fusion crusts.

Physical effects of atmospheric entry have been investigated both by numerical modeling (e.g., Love and Brownlee, 1991; Dias et al., 2016) and experimentally, in the form of static heating (e.g., Greshake et al., 1998; Toppani et al., 2001), impact ballistics (Thomas and White, 1953; Zotikov, 1959), plasma flow (Shepard et al., 1966; Loehle et al., 2017; Agrawal et al., 2018; Pratesi et al., 2018), and laser (Miley et al., 2007; Stern 2015, personal communication) set-ups. A review of the historical experiments is presented in Helber et al. (subm.). In the past, experimental conditions were generally far from those experienced by planetary material during atmospheric entry in terms of temperature, duration, and employed materials, and so far, only rarely glass could be recovered after the experiments.

In this work, petrographic and geochemical characterization of the quenched melt formed during experiments in the high-enthalpy Plasmatron facility at the von Karman Institute for Fluid Dynamics (VKI) in Sint-Genesius-Rode, Belgium, is presented. As starting material, a terrestrial basalt and an ordinary chondrite (H5) meteorite were used. The resulting glass was then compared with natural meteorite fusion crust and micrometeorites, in an attempt to advance our understanding of the thermal and chemical processes occurring during and following atmospheric entry.

2. Experimental setup, sample description, and analytical methods

The VKI Plasmatron, an induction heated plasma wind-tunnel commonly used to test spacecraft heat shields, can produce a steady-state air plasma flow of over 10,000 K temperature, and a maximum heat flux of 16 MW/m². The stagnation point region of a hypersonic body entering the atmosphere, due to the shock wave on the front of the bolide, compresses the air to temperatures exceeding 10,000 K. For this reason, the high velocity typical during atmospheric entry transforms into high total enthalpy. A subsonic ground-test facility, such as the VKI Plasmatron, is thus able to replicate the atmospheric entry flight conditions in a subsonic environment through the so-called Local Heat Transfer Simulation methodology, developed by Kolesnikov (1993). Experiments were performed in the spring of 2016 under the conditions listed in Table 1, using air as source for the plasma flow. Cylindrical samples of diameter 16 mm were cored from raw experimental materials. The sample was held in a protective, water-cooled metallic housing, maintaining a temperature range between room temperature and 70°C, before injection into the hot plasma flow. During the experiments, the material has been exposed to a cold wall heat flux of 3 MW/m² at 15 hPa as well as 1 MW/m² at 200 hPa (Table 1), corresponding to enthalpies of approximately 54 and 24 MJ/kg and flow velocities of approximately 687 and 26 m/s, respectively. These conditions reproduce the flight of an object ca. 4 m in diameter, entering with a velocity of ca. 6.5 km/s at an altitude of ca. 55 km. During the experiments, the surface temperature, the temperature of the sample in contact with the holder, and vapor ablation products at the very contact between plasma and sample were directly measured and subsequently modeled. Further detailed information on the analytical setup is provided in Helber et al. (subm.).

The specimen was placed in a hemispheric holder made of cork or graphite for different sets of experiments (Table 1). The test sample, initially at room temperature and positioned on a water-cooled holding arm, was hydraulically inserted into the plasma flow when the experimental conditions, continuously monitored, had reached the desired state. For the first set of experiments, an alkali-rich basalt was used, in order to enable the possibility of investigating evaporation phenomena of the volatile alkali metals, to act as a general analog for some basaltic achondrite meteorites and micrometeorites, and to serve as a reproducible testing material before subjecting a chondrite meteorite to full test conditions. The selected sample consists of alkali-rich basalt collected in the locality of Château de Pourcheryrolles (France), where columnar basalt occurs from the strombolian volcanic sequence of the Montpezat-sous-Bauzon, Ardèche, active between 65 Ma and 7 ka. This basalt is particularly homogeneous on the scale of a hand sample and consists of clinopyroxene phenocrysts embedded in a feldspathic groundmass, with flow fabric marked by clinopyroxene and ilmenite crystal alignment (Fig. 1a). No sulfides or sulfates were observed. According to our analyses, this alkali-rich basalt can technically be referred to as a tephry-phonolite, according to the classification by Le Bas et al. (1992). Bulk compositional data are provided in the supplemental material. Different experimental conditions were tested on the basalt, to constrain the behavior of the sample, before testing a drilled fragment of ordinary chondrite El Hammami (H5), purchased for this purpose. Ordinary chondrites represent the vast majority of all meteorites recovered on Earth (e.g., Hutchison, 2004; Grady et al., 2014). Among ordinary chondrites, type H represents a relatively reduced chemical composition and, therefore, is suitable to highlight possible redox processes induced by interaction between the melt and the atmospheric oxygen. The selected fragment contains a few relic chondrules, largely recrystallized, coarse-grained olivine (Fa₉₆₋₃₃), pyroxene (Fs₁₀₋₆₃Wo₁₋₃), and rare small plagioclase crystals and metal (Grossman, 1998).

The material recovered from experiments B 1.1 (basalt in cork sample holder) and B 1.2 (basalt in graphite sample holder; Table 1) was analyzed using a M4 Bruker Tornado μ-XRF at the Vrije Universiteit Brussel (VUB), Brussels, Belgium, for element mapping and semi-quantitative analysis. Table 1 lists the experimental conditions for the runs considered in this work.

<table>
<thead>
<tr>
<th>Experiment name</th>
<th>Material</th>
<th>Sample holder</th>
<th>Heat flux (MW/m²)</th>
<th>Heat flux probe</th>
<th>Pressure (hPa)</th>
<th>Duration (s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B1.1</td>
<td>Basalt</td>
<td>Cork</td>
<td>3.04</td>
<td>Hemispherical</td>
<td>15</td>
<td>41</td>
</tr>
<tr>
<td>B1.2</td>
<td>Basalt</td>
<td>Graphite</td>
<td>3.12</td>
<td>Hemispherical</td>
<td>15</td>
<td>21</td>
</tr>
<tr>
<td>B2.1</td>
<td>Basalt</td>
<td>Cork</td>
<td>1.01 (1.2)⁴</td>
<td>Flat (hemispherical)</td>
<td>200</td>
<td>12</td>
</tr>
<tr>
<td>GC2.1</td>
<td>Ordinary chondrite</td>
<td>Cork</td>
<td>1.01 (1.2)⁴</td>
<td>Flat (hemispherical)</td>
<td>200</td>
<td>21</td>
</tr>
</tbody>
</table>

⁴ Calculated by applying a similarity rule for the heat flux.
titative chemical analysis, with a Rh beam and spot size of 25 μm. Preliminary observations and rough chemical characterization were also performed with a JEOL JSM IT-300 scanning electron microscope (SEM), equipped with EDS detector, at the VUB. Electron microprobe analysis (EMPA) was performed with a JEOL JXA 8530 F instrument at the Natural History Museum of Vienna, Austria, equipped with five wavelength-dispersive spectrometers (WDS) and one EDS, on polished chips prepared from the material recovered from experiments B1.2, B2.1, and OCZ.1 (Table 1). Operative conditions were 15 kV acceleration voltage, 12 nA beam current, and with a fully focused beam. For glassy material, a defocused beam with a spot size of 10 μm was used. For quantification, a ZAF correction was applied and the content was calculated in oxides, except for the metal in the ordinary chondrite. Detection limits for major elements are listed in Table S1 in the Supplemental Material. The composition of olivine ([Mg,Fe]SiO₄) is expressed as fayalite Fe₂SiO₄ (Fa) mol% (Mg/(Mg + Fe)+100). The mineral abbreviations Olf for olivine and Fa for fayalite, as well as Mag for magnetite and other abbreviations, are used according to Whitney and Evans (2010). Major and trace element concentrations were additionally determined using a Teledyne CETAC Technologies Analyte G2 ArF excimer-based laser ablation system coupled to a Thermo Scientific Element XR double-focusing sector field ICP-mass spectrometer (LA-ICP-MS) at the Department of Chemistry of Ghent University, Belgium, following the procedures described in Das Gupta et al. (2017).

The oxidation state of Fe in the basalt before and after the experiments B1.1 and B1.2, as indication of the redox conditions, was determined using Fe K-edge X-ray Absorption Near Edge Structure (XANES) spectroscopy. X-ray absorption spectroscopy (XAS) spectra were collected at the LISA beamline BM08 (d’Acapito et al., 2016, 2017) at the European Synchrotron Radiation Facility (ESRF) in Grenoble (France). All samples were measured in fluorescence mode by means of a 12-element solid state (high-purity Germanium) detector. The monochromator was equipped with a pair of Si [111] flat crystals; higher order harmonics were rejected by using Si-coated focusing mirrors (Ecut = 15 keV); beam size on the sample was approximately 200 × 200 μm². The basalt was analyzed as a powder mixed with cellulose and pressed into a pellet, with an amount of material such as to keep the maximum total absorption (μ) below 1.5. From the material after experiment B1.2, XAS data were collected on molten droplets embedded in a resin pellet and then polished. Material recovered after experiment B2.1, due to the low amount of quenched melt available, required manual glass separation, and only after ascertaining the absence of crystalline phases was prepared as powder sputtered onto a kapton tape.

XAS spectra were collected along the Fe K-edge from 200 eV below the edge up to 650 eV after the edge, with variable energy step (0.10 eV across the pre-edge and 0.3 eV on the edge regions). The software ATHENA (Ravel and Newville, 2005) was used to average multiple spectra. Standard procedures (Lee et al., 1981) were followed for pre-edge background removal, spline modeling of bare atomic background, edge step normalization using a far above the edge region, and energy calibration. Pre-edge peak data analysis was done according to the method described in Wilke et al. (2001) and Giuli et al. (2002 and 2011).
3. Results

In all experiments, the plasma flow temperature was estimated to be roughly between 5000 and 9000 K (Helber et al., subm.). The temperature on the surface of the sample, measured and modeled, reached an asymptotic value in the range of 2280–2360 K, regardless of the nature of the material and of the sample holder (Helber et al., subm.). All samples started at room temperature and cooled slightly during pumping down of the test chamber to reach the required vacuum. The temperature gradient throughout the sample material strongly depended on the sample holder material (graphite or cork). In the case of graphite, which is a good thermal conductor, the temperature at the back of the sample rapidly increased by 640 K compared to the initial temperature. In the case of cork, which in contrast to graphite is a good thermal insulator, the temperature at the back of the sample almost did not vary during the experiments, with limited temperature rises differences between only 3 K and 100 K. Nevertheless, all samples exhibited melting, boiling and evaporation during the experiments. Detailed information is provided for the individual experiments.

3.1. Petrographic and geochemical observations in experiments with basalt

The response of the basalt strongly depended on the nature of the sample holder. In the case of cork, the basalt experienced intense fracturing and ejection of material, due to the insulation properties of cork, causing a strong thermal gradient within the sample between the exposed surface and the cold interior. Here, it should be noted that fracturing has been also observed after drying the sample by exposure to 100°C overnight. In the case of graphite, low-viscosity melt dropped away on the surface of the sample holder and melting also occurred along the sides of the sample, triggered by the high thermal conductivity of graphite. A few mm of sample were lost at the front of the plasma flow by vaporization and removal of melt during the experiments. Upon quenching, the melt appears as a thin (< 500 μm) film of glass coating the preserved basalt, also in ejected fragments from experiment B1.1 (Fig. 1b, c), locally splattered on the surface of the basalt (Fig. 1c, d), as droplets of blackish glass after experiment B1.2. The transition between the preserved portion of basalt and the glass is marked by abundant vesicles, with vesicles of various size (50 μm–1 mm) (Fig. 1e). Spherical objects, 200–500 μm in size, have formed at the front of the glass film (Fig. 1f, g). In the experiment B1.2, Fe-Si metal alloys crystallized at the contact between the melt and the graphite (Fig. 1h). These alloys have a calculated mineral formula of Fe2Si, which is in between that of hapkeite (Fe-Si) and gupeite (Fe-Si), both of extraterrestrial origin and observed in lunar soils and likely in metallic cosmic spherules, respectively. Hapkeite has been suggested to form by condensation of impact vaporized phases (Anand et al., 2003). In BSE-SEM images, local schlieren and domains with different BSE contrast in BSE-SEM images can be observed (Fig. 1h). Locally, evidence of devitrification and incipient crystallization, with skeletal growth of unidentified phases over relic nuclei, is present (Fig. 1i).

Chemically, the glass formed from the basalt shows overall depletion in volatile alkalali metals (40–60% of the original coating on Na, 50% for K) and highly volatile elements, with apparent enrichment in more refractory elements (Mg and Ca contents are doubled; Fig. 2a, Tables S2–3). In detail, the difference between the bright and dark glass schlieren in BSE images is limited to a more marked enrichment in Ti, Fe, and Mn in the bright domains than in those with darker appearance. The enrichment in Ti, Ca, Mg, Fe, and Mn with a factor 2 or more in the glass is common, regardless of the sample holder. Additionally, in experiment B1.2, a considerable enrichment in Ni (20 times higher than the preserved basalt) and Cr in the glass is observed. Spectroscopic analysis during the experiments indicates that Na, K, Mg, Ca, Fe, and Ti are volatilized, but the relative abundance could not be unambiguously determined, because both the cork and the graphite produced a strong K and Na emission in addition to those from the samples (Helber et al., subm.). Relative to the pristine basalt before the experiments, the glass is enriched in rare earth elements (REE) and Y (Fig. 2b) and in moderately siderophile elements, such as Co, Ni, and Cu (Table S5). Among the spherical objects, two populations can be distinguished for their size. While the observed larger objects (Fig. 1f, g) have a clear, relatively “feldsparitic” composition (Table S2), the composition of smaller objects was difficult to be determined, due to contamination from the surrounding unmelted basalt. According to the pre-edge peak analysis of XAS data (reported in Fig. 3), the glass in both experiments B1.2 and B2.1 (0.17 ± 0.05 and 0.11 ± 0.05, respectively) exhibits a lower Fe3+/Fe2+ ratio than that in the original basalt (0.75 ± 0.15).

3.2. Petrographic and geochemical observations in the experiment with the ordinary chondrite

The meteorite sample experienced only melting and limited evaporation during the experiment. No material was ejected or escaped from the sample holder. The resulting glass displays large vesicles and coats the exposed surface of the sample (Fig. 4a). There is no sharp transition between the unaffected interior of the meteorite sample and the molten and quenched portion and the glass is largely crystallized. Olivine crystals in contact with the glass exhibit cracks filled by glassy melt, trails of metal-rich inclusions, and an overgrown rim with a higher Mg/Fe ratio than that of the host olivine (relative average Mg# from 81 to 85; Fig. 4b, Table S4). The glass resulting from the
siderophile elements (Co, Ni, Cu) are highly variable and difficult to interpret, likely due to the large internal variability in the ordinary chondrite data (Table S5) and to the limited laser beam diameter during LA analysis (50 μm) in comparison to the coarse grain crystal size in the ordinary chondrite (e.g., Fig. 4).

Olivine that crystallizes from the melt can be subdivided into two populations on the basis of the MgO content with respect to the olivine in the interior of the meteorite (Fig. 5b). However, both populations exhibit similar enrichment in refractory elements, such as (Al, Cr, Ni, Ca, but not Ti) with apparent depletion in the moderately volatile elements and Mn. In particular, newly crystallized olivine seems to contain “impurities”, such as Cr, Ni, and Ca, even though this might be apparent and due to analytical limitations. In fact, the apparent enrichment in Na and Al is likely related to local contamination from the groundmass, due to the size of the analyzed material (olivine rims have a thickness of a few μm; Fig. 4) with respect to the interaction volume of the electron microprobe beam (also on the order of some μm; Table S4).

4. Discussion

4.1. Melting, evaporation, and condensation

In all experiments, regardless of the nature of the material and of the sample holder, the resulting glass composition clearly shows minor enrichment in some moderately refractory elements (e.g., Ti and Ca) and relative depletion in highly volatile elements (such as K and Na). The evaporation of highly volatile alkali metals is particularly obvious in the case of the basalt, as the selected basalt has significantly higher concentrations of Na₂O (6.7 wt%) and K₂O (5.8 wt%; Table S2) relative to the H chondrite (Na₂O of 1.6 wt% and K₂O of 0.15 wt%; Table S6). The formation of vesicles in both the basaltic and chondritic glasses is obviously related to the vaporization of such highly volatile elements (50% condensation temperature Tc of 958 K and 1006 K for Na and K, respectively; Lodders, 2003) with a possible contribution from elements such as Fe (50% Tc of 1334 K; Lodders, 2003). The latter element was detected by spectroscopic analyses and is observed as Fe-oxide condensates along the rims of the vesicles. Vesculation is commonly observed in natural meteorite fusion crusts (e.g., Genge and Grady, 1999a; Pittarello et al., 2019), but also in micrometeorites (e.g., Genge, 2016b; Suttle et al., 2019), where a high content of volatiles in the original precursor material has been invoked to explain vesicle formation. In the case of the basalt, a high abundance of volatiles, such as water and halogens, is likely present. However, this is not the case for the ordinary chondrite, which in contrast to carbonaceous chondrites is relatively dry (e.g., Brearley, 2006). The quenched melt in the experiments does not display any lateral variation reflecting melting of local phases but is rather homogeneous. This observation is consistent with what observed in natural fusion crusts of H5 chondrites (Pittarello et al., 2019) and in contrast with previous works (e.g., Genge and Grady, 1999a; Thaiss and Taylor, 2009). However, in the case of the experiments, the small sample size and the geometry of the experiments favor mixing and homogenization. In natural fusion crusts, homogenization may be additionally aided by the spinning of the meteorite during flight.

The origin of the spherical objects found at the interface of the glass formed from the basalt remains unclear. In some cases, chemical compositions rich in alkali metals and silica might suggest possible condensation of vaporized phases. No spherical objects were observed in the glass obtained from the ordinary chondrite. In natural fusion crust, a possible ablation sphere, with a composition close to that of the bulk chondrite, was observed (Pittarello et al., 2019). This suggests that the observed spherical objects on the surface of the basaltic glass formed by a different mechanism than meteorite ablation spheres.
Moderately siderophile elements, such as Co, Ni, and Cu are enriched in the glass formed in all the experiments. Partitioning of such elements between melt and silicates may be controlled by the oxygen fugacity (Gaetani and Grove, 1997). In this case, the partitioning would be between vapor and melt. The enrichment in moderately siderophile elements shown in the glass is likely the result of the evaporation of the more volatile elements. However, despite the relatively large uncertainties associated with these analyses, simple mass balance calculations suggest passive enrichment not to be the sole cause for the observed concentration ranges.

4.2. Redox processes

For the selected basalt, Log $f(O_2)$ was estimated with the software MAGMA (Schaefer and Fegley, 2004), yielding $-$4 at 2200 K, which is close to the temperature reached during the experiments. Oxygen partial pressure in the gas during the experiment can be calculated. Such estimates suggest high $f(O_2)$, such as Log $f(O_2) = 1.25$ and Log $f(O_2) = 1.41$ assuming chemical equilibrium, which represents a relatively oxidizing environment during the experiments. This is valid if we ignore the potential reducing effect induced by the presence of graphite and the organic matter in the sample holder, although the geometry of the experiment ensures that any interaction between the melt and the sample holder is generally avoided, except at the very contact between the two. This local effect is negligible in the scope of the overall redox reactions. The $Fe^{3+}/Fe^{tot}$ ratios observed in the experiment B1.2 with graphite sample holder ($Fe^{3+}/Fe^{tot} = 0.17 \pm 0.05$) and in experiment B2.1 with cork sample holder ($Fe^{3+}/Fe^{tot} = 0.11 \pm 0.05$) are very close to those calculated according to the Kress and Carmichael (1991) model using the glass composition, the estimated T and oxygen fugacity conditions of the experiment. This correspondence suggests that the melt was in equilibrium (or close to equilibrium) with the surrounding atmosphere. Despite the short duration of the experiments, it is not surprising that equilibrium can be achieved for the Fe redox reaction, because at extremely high temperature, as recorded during the experiments, Fe redox kinetics are extremely fast. In an experiment, where basalt was molten at ca. 1770 K, equilibrium was reached after 66 s (Gicconi et al., 2015). The experimental data indicate that at such high temperature the produced melt can quickly equilibrate with the surrounding medium and, for the conditions used here, produce a melt where $Fe^{3+}$ and $Fe^{5+}$ coexist (in our case with the $Fe^{3+}/Fe^{tot}$ ratio ranging between 0.11 and 0.17). These data are compatible with the formation of magnetite in the produced chondrite melt, where the oxygen fugacity in H chondrites has been calculated to be ca. Log $f(O_2) = 20$ at 800 °C (Rubin et al., 1988). It must be noted that, as the melt achieved equilibrium, the resulting $Fe^{3+}/Fe^{tot}$ ratio would have been the same regardless if the starting material was more oxidized (as in the basalt case) or more reduced (as in the meteorite case).

![Fig. 4](https://example.com/image.png)

Fig. 4. Melt formed in the experiment OC2.1, from a fragment of the H5 ordinary chondrite El Hammani, and comparison with natural meteorite fusion crust. a) Microphotograph of the thin chip used for analysis, showing the interior of the meteorite and the vesiculated melt. Reflected light, optical image. b) BSE-SEM image of the transition zone between the unaffected interior of the meteorite and the melt, showing olivine coroned by cracks and by tails of opaque phases and with a darker rim towards an apparently homogeneous melt. c) BSE-SEM image of olivine relic clast exhibiting overgrowth from the melt with initial crystallization of Mg-rich olivine, followed by crystallization of progressively more Fe-rich olivine. Note the fine grained skeletal phases crystallized in the melt. d) BSE-SEM image of hopper and skeletal olivine crystallizing from the melt. e) BSE-SEM image of the groundmass between new crystals of olivine, with skeletal magnetite and likely olivine. f) BSE-SEM image of a metal bleb in the melt consisting of botryoidal kamacite in a trolledite matrix. g) BSE-SEM image of a chromite crystal in a groundmass consisting of likely skeletal olivine, trolledite/blebs, and glass. h) BSE-SEM image of zoning in olivine clasts and crystals in the Antarctic meteorite Asuka 090502, classified as H5 (Pittarello et al., in review). Note the similarity with (c). i) BSE-SEM image of hopper olivine with skeletal magnetite in the groundmass in the Antarctic meteorite Asuka A 09004, classified as H5 (Pittarello et al., in review). Note the similarity with (c).
Experiments with sample holders of different materials, namely graphite and cork, were performed to control the possible influence of sample holders on the resulting glass. While graphite is a good heat conductor, cork is a good insulator. Chemically, both contain carbon, a potentially reducing agent, but because of their individual thermal response, carbon is released under different conditions. The thermal properties of the two materials affected the temperature rise of the samples at the exposed surfaces (Table 1), as the basalt embedded in graphite reached boiling temperatures much earlier than that inserted in cork. At a local scale, at the direct contact between the quenched melt and the graphite, metal droplets (Fe-Si alloys) crystallized (Fig. 1h), suggesting that the interaction between the still hot melt and graphite induced chemical reduction of the melt. However, considering the geometry of the experimental set up, the glass not in direct contact with the graphite holder should not have been chemically affected by the sample holder.

The redox process is more complex in the case of the ordinary chondrite, which was originally relatively reduced (e.g., Rubin et al., 1988), and contains metallic Fe and Ni. Nevertheless, considering olivine crystallization as a proxy for the redox process occurring in the melt, an initial reduction is indicated by the occurrence of olivine compositions at the direct contact with relict olivine crystals with higher Mg/Fe ratios than those of olivine crystals in the interior of the meteorite. However, this observation might reflect the melt temperature, as Mg-rich terms have higher crystallization temperature than the Fe-rich terms. Nevertheless, in experiment OC2.1, oxidation is the dominant process, as suggested by the crystallization of magnetite in the groundmass, the absence of taenite in the melt, and the growth of Fe-rich olivine at the outer rim of relict olivine fragments in the melt or as dendrites (Fig. 4). Even though no significant variations in the amount of magnetite have been observed in this experiment, previous works have related the abundance of magnetite in the groundmass to the extension of oxidation, suggesting that oxidation shows a gradient from the interior of the meteorite to the fusion crust (e.g., Genge and Grady, 1999a, 1999b). Additionally, we have identified <5 μm skeletal chromite grains in the quench melt of our ordinary chondrite sample. Chromite is essentially absent in low grade type-3 ordinary chondrites, but grew during metamorphism in the ordinary chondrite parent body/ies and is present as relatively large (−50–100 μm) anhedral grains in higher grade specimens such as our test sample (Ramdohr, 1967; Huss et al., 1981; Rubin, 2003; Wlotzka, 2005). Although it is possible that the grains we have found may represent surviving relics, their small size and skeletal structural seem to imply fast crystallization from the experimentally produced melt (albeit possibly over relict chromite nuclei). Such chromite grains may indicate further oxidation of Cr₂⁺ originally hosted in silicate minerals to Cr³⁺ and the stabilization of chromite in the newly formed melt.

In conclusion, our observations that (i) the basaltic glass is generally “reduced” with respect to the highly oxidized pristine basalt and that (ii) magnetite and Fe-rich olivine rims occurs in the chondritic glass, indicating oxidation of the originally relatively reduced ordinary chondrite, all point to an equilibrium redox condition proper of the experimental setup. This equilibrium condition is rapidly reached at high temperature, within the duration of the experiment. The characteristic oxidation state of the experimental setup is intermediate between that of the basalt and that of the chondrite. Disregarding the original oxidation state of the material, the equilibrium depends on temperature, composition and oxygen fugacity, and it is reached by varying of the relative amounts of Fe³⁺ (in the chondrite), Fe²⁺, and Fe³⁺.

4.3. Comparison with meteorite fusion crust, ablation spherules, and melted micrometeorites

The experiment OC2.1 aimed to reproduce meteorite fusion crust by simulating atmospheric entry of a fragment of ordinary chondrite. The quenched melt produced in the experiment presents the same features observed in natural meteorite fusion crusts from chondrites (for a detailed description, see Genge and Grady, 1999a; and Pittarello et al., 2019), such as extensive crystallization of olivine, the presence of magnetite in the groundmass, and the peculiar zoning in the overgrown olivine (Fig. 3). The extensive crystallization of the melt after experiment OC2.1 was not expected, considering the contrasting conditions with respect to the natural atmospheric entry, such as the duration of the experiment (seconds vs. minutes), and the size of the sample (14 mm in diameter of exposed surface). However, the cooling rate seems to have been similar to that experienced by meteoroids, allowing crystallization of olivine. Thus, based on a comparison of the resulting material with the natural samples, the experiment successfully reproduced the (chemical) conditions encountered by asteroidal material when entering Earth’s atmosphere.

Micrometeoroids experience partial (similarly to regular meteoroids) to complete melting (i.e., cosmic spherules), involving evaporation of (moderately) volatile components, oxidation of Fe-Ni alloys to Fe oxides (such as magnetite or wüstite, whose presence has been used to determine the oxygen fugacity), reduction of Fe and formation of immiscible metal beads when the precursor material is particularly enriched in carbon, and loss of siderophile and potentially chalcophile elements as a result of Fe-Ni metal and sulfide bead mechanical ejection or spinel group mineral expulsion (e.g., Taylor et al., 2005; Cordier et al., 2011; Genge, 2016a). Evaporation in cosmic spherules can reach up to 60%–90% in the Ca-Al-Ti-rich (therefore, more refractory) terms, depending on the original composition (Herzog et al., 1999; Alexander et al., 2002; Engrand et al., 2005; Taylor et al., 2005; Cordier et al., 2011), however, condensation products have never been identified in micrometeorites. More recent studies have highlighted that the pres-
ence of volatile elements in micrometeorites might be responsible for the fracturing and heating of particles during atmospheric entry, favoring interaction with atmospheric gases (e.g., Suttle et al., 2019).

The most important process controlling the melt evolution during the atmospheric entry is the oxidation of Fe. This mostly implies the formation of magnetite crystals and, locally, the formation of new, relatively FeO-rich olivine, generally overgrowing pre-existing olivine crystal nuclei (Genge et al., 1997; Genge and Grady, 1999a, 1999b; van Ginneken et al., 2017), similarly to what is observed in meteorite fusion crust and in the experiment OC2.1. However, reduction processes may play an important role during the atmospheric entry of micrometeoroids when the precursor material is particularly rich in carbon (e.g., Genge and Grady, 1999a). This leads to the formation of carbon and metal-rich liquids that are immiscible with the surrounding silicate melt and tend to be ejected from the host material (i.e. pyrolysis; Genge and Grady, 1998).

The maximum temperature achieved during atmospheric entry of micrometeoroids might have been different from that reached in the experiments (ca. 2200 K in the experiments vs. likely 900–2000 K for micrometeoroids; e.g., Love and Brownlee, 1991; Taylor et al., 2000), as well as the oxygen fugacity related to the altitude, where melting can occur for the specific material. Additionally, micrometeorites experience peak heating at much higher altitudes in the atmosphere, where the partial pressure of oxygen is significantly lower (e.g., Genge, 2016a; Genge et al., 2017). This implies that a direct comparison between the results of the experiments presented here and the features contained in micrometeorites might not be completely appropriate. Further experiments, using several precursor materials (including carbonaceous and iron meteorites) are planned in the attempt to reproduce the variety of textural, chemical and isotopic features observed in micrometeorites.

5. Conclusions

The quenched melt recovered from a series of melting experiments in a plasma facility, involving both basaltic and ordinary chondritic starting materials, provides constraints on the thermal and chemical processes that affect extraterrestrial material during atmospheric entry. Chemically, the composition of the resulting glass is depleted in highly volatile alkali metals and slightly enriched in moderately siderophile elements. In the experiments involving the alkali basalt, spherules derived from the condensation of the volatilized elements were observed. In terms of redox reaction, the selected experimental conditions affect the redox state of the samples. Regardless of the initial conditions of the starting material, equilibrium between the sample and the surrounding medium is rapidly reached. For this reason, the terrestrial basalt (with a starting Fe\(^{3+}\)/Fe\(^{2+}\) ratio of 0.75) has been reduced to a Fe\(^{3+}\)/Fe\(^{2+}\) ratio of 0.11–0.17 ± 0.05, within the stability field of olivine + magnetite. In contrast, the chondrite is oxidized, as suggested by the crystallization of magnetite, which indicates the presence of Fe\(^{3+}\). However, the path to reach this equilibrium might have not been linear, with possible reduction during the initial stages of melting in the ordinary chondrite. In general, the microstructures observed in the material recovered after experiment OC2.1, with an HS ordinary chondrite, compare well to whose present in melted micrometeorites and in natural meteorite fusion crusts from the same type of chondrite. As the experiments have demonstrated to have successfully reproduced the features observed in natural samples, the thermal and redox conditions likely encountered by meteoroids during atmospheric entry have been closely approximated by our experiments.

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Appendix A. Supplementary data

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References


