# The Stubenberg meteorite-An LL6 chondrite fragmental breccia recovered soon after precise prediction of the strewn field

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#### Abstract :

On March 6, 2016 at 21: 36: 51 UT, extended areas of Upper Austria, Bavaria (Germany) and the southwestern part of the Czech Republic were illuminated by a very bright bolide. This bolide was recorded by instruments in the Czech part of the European Fireball Network and it enabled complex and precise description of this event including prediction of the impact area. So far six meteorites totaling 1473 g have been found in the predicted area. The first pieces were recovered on March 12, 2016 on a field close to the village of Stubenberg (Bavaria). Stubenberg is a weakly shocked (S3) fragmental breccia consisting of abundant highly recrystallized rock fragments embedded in a clastic matrix. The texture, the large grain size of plagioclase, and the homogeneous compositions of olivine (Fa(31.4)) and pyroxene

(Fs(25.4)) clearly indicate that Stubenberg is an LL6 chondrite breccia. This is consistent with the data on O, Ti, and Cr isotopes. Stubenberg does not contain solar wind-implanted noble gases. Data on the bulk chemistry, IR spectroscopy, cosmogenic nuclides, and organic components also indicate similarities to other metamorphosed LL chondrites. Noble gas studies reveal that the meteorite has a cosmic ray exposure (CRE) age of 36 +/- 3 Ma and that most of the cosmogenic gases were produced in a meteoroid with a radius of at least 35 cm. This is larger than the size of the meteoroid which entered the Earth's atmosphere, which is constrained to <20 cm from short-lived radionuclide data. In combination, this might suggest a complex exposure history for Stubenberg.

#### **INTRODUCTION**

On Sunday March 6, 2016 at 21:36:51 UT (corresponding to the late local evening, UT+1h), extended areas of Upper Austria, Bavaria, and the southwestern part of the Czech Republic were illuminated by a very bright fireball. In particular, the inhabitants of Bavaria, where the skies were mostly clear, were stunned by the luminous and sonic effects of the fireball. Observation was difficult in most of Central Europe due to a cloudy sky. Fortunately, parts of western Czech Republic also had partly clear skies in some places and it enabled researchers to obtain the instrumental records provided by the digital autonomous observatories (DAFOs) of the Czech part of the European Fireball Network. These photographic and radiometric records proved to be essential to describe this rare natural event in detail. The cameras at six Czech stations photographed the event at least partly and almost all Czech cameras recorded its light curve by fast photometers with high time resolution of 5000 samples per second. Thanks to these records, the fireball could be described rigorously and with a high reliability and precision (Spurný et al., 2016). Moreover, the impact area of potential meteorites could also be computed and the ground distribution of meteorites could be modeled. This prediction was very precise, because all six recovered meteorites were found almost exactly in the predicted location for the given meteorite masses (Spurný et al., 2016).

In this paper the mineralogical and chemical characteristics of this new meteorite will be presented including data on organics, short-lived cosmogenic radioisotopes, and noble gases. The meteorite "Stubenberg" has been accepted by the Nomenclature Committee (The Meteoritical Bulletin, 2016, in preparation) and preliminary data were published by Ebert and Bischoff (2016) and Spurný et al. (2016). After the meteorite falls of Braunschweig (L6; Bartoschewitz et al., 2017), Žd'ár nad Sázavou (Spurný et al., 2016), and Ejby (H5/6; The Meteoritical Bulletin, 2016, in preparation), Stubenberg is the fourth recovered meteorite fall within less than three years impacting Earth only ~800 km apart. Less than 10 meteorite falls/year are recovered globally, and hence, this concentration of recorded falls is remarkable.

## SAMPLES AND METHODS - SUMMARY

In order to obtain information about the classification of the rock and to reveal details on chemical, physical, and mineralogical features and characteristics of the rock different analytical methods were applied. This is a brief summary and further details are given in the "Supplementary Materials". Several thin and thick sections and fragments of Stubenberg were

studied using optical and electron microscopy. Electron microscopic studies with various Scanning and Transmission Electron Microscopes (SEM, TEM) were performed in Münster, Jena, Prague, and Munich and chemical data by electron microprobe were obtained in Münster. Raman spectroscopical investigations (Munich) were conducted with a Horiba Xplora Integrated confocal Laser micro Raman system on an Olympus BX51 microscope.

After homogenizing a 0.5 g-piece, a 200 mg whole-rock sample of fragments and splinters was analyzed for major and trace element concentrations by inductively coupled plasma atomic emission spectroscopy (ICP-AES) and inductively coupled plasma sector field mass spectrometry (ICP-SFMS) following the procedure described by Barrat et al. (2012, 2015, 2016). Cosmogenic radionuclide concentrations were analyzed by means of non-destructive high purity germanium (HPGe) gamma spectroscopy in the underground laboratories at the Laboratori Nazionali del Gran Sasso and at the Max-Planck-Institut für Kernphysik in Heidelberg (Arpesella, 1996, Heusser et al., 2015), and bulk oxygen isotope composition was analyzed by means of laser fluorination in combination with gas source mass spectrometry (Sharp, 1990). The data were normalized to the composition of San Carlos olivine ( $\delta^{17}O_{VSMOW2} = 2.715 \%$ ,  $\delta^{18}O_{VSMOW2} = 5.220 \%$ ; see Pack et al., 2016).

For the Ti-Cr isotope analyses sample powders of Stubenberg (2 x 50 mg) were dissolved and purified using ion exchange chromatographic procedures. Titanium isotope analyses were then carried out using a Thermo Fisher Scientific Neptune Plus multicollector-inductively coupled plasma mass spectrometer (MC-ICPMS) both at the Institut für Planetologie in Münster and at the Institute of Geochemistry and Petrology at ETH Zurich, while Cr isotopes were analyzed in Zurich only. The analytical procedures followed those described in Schönbächler et al. (2004), Williams (2015), and Zhang et al. (2011) for Ti and in Steele and Schönbächler (2016) for Cr and are described in more detail in the "Supplementary Materials". The noble gases He-Xe present in two bulk samples "I" (23.3 mg) and "II" (71.5 mg) were analyzed at ETH Zurich using the standard procedures described in detail by Riebe et al. (2017, in revision).

Regarding physical properties, Fourier transform infrared spectroscopy (FTIR) spectroscopic measurements were undertaken using a Bruker Vertex 70v at the IR/IS laboratory located at the Institut für Planetologie (Münster) and the Magnetic Susceptibility (MS) of various Stubenberg fragments was measured in the strewn field as well as in Nördlingen with the SM30 of ZH Instruments. Additional instrumental details are given in the "Supplementary Materials". A destructive methanolic solvent extraction of organic matter was performed in an

agate mortar using 50 mg freshly broken fragment of the Stubenberg meteorite. Analytical details and information on the data processing are given in Schmitt-Kopplin et al. (2010, 2012), Tziotis et al. (2011), Hertkorn et al. (2015), and Ruf et al. (2017).

# RESULTS

## Fall circumstances and geographic details of find area

From the detailed light curve of the observed bolide taken by the fast photometers (time resolution 5000 samples/s) as part of the Czech fireball automated cameras (Spurný et al. 2007), it was evident that the initial meteoroid heavily fragmented in the last third of its luminous atmospheric flight. A very complex model of the impact area was produced using precise trajectory data combined with the known wind profile available from balloon data. The distribution pattern of individual meteorites deposited from each particular breakup were directly visible on the light curve, which will be described along with the bolide data elsewhere. The resulting predicted impact area near the village of Stubenberg in Lower Bavaria has a complicated irregular shape (Fig. 1). This prediction was convincingly confirmed by the positions and masses of the recovered meteorites, as described in Fig. 1. The northern part of the strewn field is dominated by forest at an elevation of about 500 m above sea level, in which the samples M5 and M6 were found (Table 1). Most other areas represent farmland (Fig. 1). Stubenberg is a small village in Lower Bavaria with a population of about 1400 residents.

#### **Petrography and Mineralogy**

#### Description of hand specimens

All recovered stones of the Stubenberg meteorite are shown in Fig. 2 and listed in Table 1. The first fragments of one broken individual stone were found six days after the fall. Within several weeks of intensive search six meteorite individuals with a total mass of 1473 g were recovered. The largest fragment (M5) has a mass of 1320 g (Figs. 1 and 2e; Table 1), and a bulk density of  $3.29 \pm 0.01$  g/cm<sup>3</sup>. The first recovered piece was heavily-fragmented at the impact location, when hitting the ground. The main mass broke into 3 larger pieces and several smaller fragments (M1a - M1n; Table 1) covering an area of ~10 meters in diameter. From the time of impact until recovery the fragmented piece remained in place. The biggest fragment M1b has a weight of 23.58 g (Fig. 2a); all pieces a 0.1 to 0.3 mm thick, dark-brown

to black and fusion crust is mostly well preserved. While the surface of the meteorite is predominantly smooth, the fusion crust shows polygonal cracks in several spots. Meteorite piece M6, weighing 35.89 gram (Fig. 2f), was found as a fragmented individual with two almost perpendicular fracture surfaces not covered by fusion crust. Intensive search did not yield the missing fragments. Close inspection by SEM later showed abundant FeS melt splashes and rare glass droplets on the broken surfaces, suggesting a high altitude break-up event, possibly at the end or slightly after the luminous flight. Cutting the meteorite by wire saw revealed a rather friable rock.

Internally, a greyish, silicate-rich component dominates the meteorite, rare mm-sized fragments are visible, and minor amounts of minerals with metallic lustre are recognizable throughout the rock. Rarely, shock veins are observed consisting of the metallic phases and troilite as confirmed by detailed mineralogical studies on thin sections. Since the rock was found about one week after the fall, it has already been affected by some minor weathering processes as documented in places by the brownish taint on the surfaces of the broken piece.

## Mineralogy – texture and compositions of phases

The brecciated meteorite has abundant fragments with a recrystallized texture and a small number of chondrule relics (Fig. 3). Especially the outlines of former barred-olivine (BO) and radial pyroxene (RP) chondrules are still recognizable (Fig. 3). One of these relic chondrules is a several mm-sized macrochondrule (Weyrauch and Bischoff, 2012). The distribution and obtained abundances of various phases are shown in Figures S1 and listed in Table S1.

Olivine is by far the most abundant phase (53-56 vol%; Table S1). Generally, the grains are variable in size and show strong undulatory extinction. The olivines are homogeneous in composition throughout all thin sections. The mean composition of 54 analyzed olivines is Fa<sub>31.4±0.3</sub> with a compositional range between 30.5 and 32.2 mol% Fa (Table 2). The low-Ca pyroxenes (~17 vol%) and Ca-pyroxenes (6.3 vol%) have mean compositions of Fs<sub>25.4</sub>Wo<sub>2.0</sub> and Fs<sub>10.3</sub>Wo<sub>43.9</sub>, respectively (Table 2; Fig. S1). The 43 analyzed low-Ca pyroxenes vary between 24.8 and 26.1 mol% Fs. Ca-pyroxene has a much lower modal abundance compared with low-Ca pyroxene and is also homogeneous in composition. Raman mapping supports that the dominating phases of Stubenberg are olivine, orthopyroxene, Ca-pyroxene, and Abrich plagioclase. According to the Raman spectra all pyroxenes show very uniform compositions with insignificant variations not only in one single clast, but also from clast to clast. Selected olivine and pyroxene particles have been tested for water (H<sub>2</sub>O/OH) contents

by Raman spectroscopy, but no Raman signals could be detected in the  $H_2O$  Raman spectral region (3400-3700 cm<sup>-1</sup>) due to possible rapid terrestrial weathering of the mineral constituents at the impact location.

Plagioclase (10-13 vol%; Table S2) is present as quite large crystals often exceeding 100  $\mu$ m in size (Fig. 3). Mean plagioclase has An- and Or-components of  $11.1 \pm 0.4$  and  $5.5 \pm 1.2$  mol%, respectively (n= 42; Table 2). The compositions vary between 10.3 and 12.1 mol% An.Both the phosphates merrillite and Cl-apatite were found in the studied thin sections. Cl-apatite is more abundant than merrillite. Their compositions are listed in Table 2.

Metal and sulfide occur as small grains distributed throughout the entire thin sections. Metallic phases are homogeneously distributed and up to 250  $\mu$ m in apparent size. The chemical compositions of these phases are given in Table S2. In general, kamacite has mean Ni- and Co-concentrations of 3.9 and 6.1 wt%, respectively. The taenite composition is variable, with Ni content varying from 41.9 to 48.2 wt% (average ~44.3 wt% Ni, ~1.8 wt% Co, 0.22 wt% Cu; n= 38). Some grains may be tetrataenite due to slight anisotropy in reflected, polarized light.

Troilite is an abundant phase in Stubenberg (approx. 9 vol%; Table S1) and is distributed throughout all thin sections, occasionally forming mm-sized aggregates. An approx. 1.5 mm long, string-shaped aggregate of troilite and Ni-poor pentlandite (molar Fe/Ni ~3.3; Table S2) was observed (Fig. 4). Small amounts of pentlandite also occur in association with other troilite grains. Pentlandite was confirmed by TEM using selected area electron diffraction (SAED) obtained on a FIB section (Fig. 4c). At the TEM scale the pentlandite shows strong mosaicism, i.e. poor crystal quality, and sharp boundaries to neighboring troilite. {111} and {0-11} of pentlandite are nearly parallel with (001) and {010} of troilite, respectively (Fig. 4d). This is close to the orientation relationship observed for troilite/pyrrhotite and exsolved pentlandite in terrestrial and meteoritic samples (Harries and Langenhorst, 2013). The coexisting troilite shows good crystal quality and contains abundant subgrain boundaries. Superstructure reflections of pyrrhotite are absent in SAED patterns, indicating nearly stoichiometric FeS.

Chromite occurs as small grains and has mean MgO-,  $Al_2O_3$ -, and TiO<sub>2</sub>-concentrations of 1.5 wt%, 5.2 wt%, and 3.4 wt%, respectively (n= 14; Table 2). Chromite has been frequently found in all fragments. It shows anhedral grains up to 100 µm and is, according to its Raman spectrum, a high iron low magnesium member of the solid solution. Rarely some grains show deviating peak positions, which point to possibly higher concentrations of Mg and/or Al.

Further accessory phases observed are ilmenite with grain sizes up to 50  $\mu$ m (associated with taenite and troilite, or with chromite), metallic copper (grains <10  $\mu$ m, associated with kamacite, taenite), rutile, anatase, and carbon phases (graphite to heavily disordered graphite). The latter three phases were identified by Raman spectroscopy. The two different TiO<sub>2</sub>-phases anatase and rutile could be clearly identified as single grains by their different Raman spectra (Fig. S2). Rutile displays white to colourless grains up to 20  $\mu$ m. Both phases are found in direct contact with ilmenite and chromite. Carbon phases have mainly been detected decorating grain boundaries between silicate and sulfide grains. According to the band width of the main Raman bands they vary between well crystallized graphite and heavily disordered phases. As all measurements have been conducted on simply broken surfaces the disorder cannot be an artifact due to mechanical treatment and contamination by grinding or coating can be excluded. On the broken surface large grains of olivine, pyroxene, feldspar, and troilite are often recognizable as well as 120° triple junctions between olivine, pyroxene and plagioclase (Fig. 5).

# Shock metamorphism and weathering.

As stated above, based on the study of the thin sections Stubenberg is brecciated (Fig. 3a). The thin sections only show highly-recrystallized fragments having only very rare relic chondrules (type 6). Several shock veins were observed (Fig. 3d). Plagioclase and olivine show undulatory extinction, and the olivines show distinct sets of planar fractures (Fig. 3f). High pressure phases (such as ringwoodite) were searched for by Raman spectroscopy, but could not be detected even in the melt veins. No glass-like or recrystallized plagioclase phases could be found (including feldspar-glasses or maskelynite). The optical features indicate that the rock is weakly shocked (S3; Stöffler et al., 1991; Bischoff and Stöffler, 1992).

The residence time of about 7 days on the ground resulted in first signs of weathering. Locally, areas with a brownish taint are already present resulting from the first breakdown of metals. Though a few rusty spots are clearly recognized with the naked eye on the surface of all samples, alteration phases such as iron oxides-hydroxides or iddingsite (olivine alteration) could not be detected by Raman spectroscopy.

# **Chemical Characteristics**

Bulk Chemistry

A number of small interior fragments of about 0.5 g free of fusion crust were provided for chemical characterization. The results summarized in Table 3 and shown in Fig. 6 indicate that the chemical composition of Stubenberg is close to the composition of Chelyabinsk (Righter et al., 2015) and that the Stubenberg chondrite is not chemically anomalous.

# Noble gases

Noble gas results for the two interior fragments of Stubenberg sample M1h (described here as Stubenberg I and II) are summarized in Table 4. All noble gases can be explained with a mixture of cosmogenic, radiogenic, and a very small amount of heavy (Ar-Xe) trapped noble gases. Most importantly, Stubenberg does not contain solar wind noble gases (Fig. 7), because of the low <sup>4</sup>He and <sup>20</sup>Ne concentrations that are entirely of cosmogenic (cos) and radiogenic (rad) origin, the lack of trapped <sup>20</sup>Ne, and the very small concentration of trapped (tr) <sup>36</sup>Ar (Table 4). Hence, Stubenberg cannot be a regolith breccia.

The heavier noble gases are all meteoritic, as shown with the (low) heavy element ratios  ${}^{36}\text{Ar}_{tr}/{}^{132}\text{Xe}$  (=20 and 38, respectively) and  ${}^{84}\text{Kr}/{}^{132}\text{Xe}$  (=0.78 and 0.74), and the high  ${}^{40}\text{Ar}/{}^{36}\text{Ar}$  (=3900 and 4100) and  ${}^{129}\text{Xe}/{}^{132}\text{Xe}$  (=1.90 and 1.95) ratios.

We used the depth- and chemistry-dependent model by Leya and Masarik (2009) to determine the cosmic ray exposure (CRE) age of Stubenberg, assuming a simple " $4\pi$ " irradiation in space and taking into account the bulk chemistry for Stubenberg given in this work (Table 3). Using the LL chemistry that is given in the Leya and Masarik (2009) data tables yields the same results. We did not re-normalize the radii according to density. All radii in the Leya and Masarik work are given for meteoroids with a density of 3.50 g/cm<sup>3</sup>. The typical LL meteorite density might be 3.29 g/cm<sup>3</sup> (Wilkison and Robinson, 2000), which would increase all radii by 6 %. However, we do not exactly know the bulk density of the Stubenberg meteoroid, i.e. cannot take into account possible macroporosity.

The measured <sup>3</sup>He and <sup>21</sup>Ne concentrations are entirely cosmogenic, the <sup>38</sup>Ar is corrected for 2 and 4 % trapped <sup>38</sup>Ar, respectively (see Table 4 for details). As Ne is entirely cosmogenic, the measured <sup>22</sup>Ne/<sup>21</sup>Ne ratio can be used directly as a shielding indicator. The precisely measured <sup>22</sup>Ne/<sup>21</sup>Ne ratio in Stubenberg I of 1.0826±0.0026 suggests a pre-atmospheric radius of  $\geq$  35 cm. The pre-atmospheric mass of 600 kg, based on fireball observations (Spurný et al., 2016), yields the same radius of 35 cm. A higher mass within a conservative factor of two (1200 kg; radius of ~45 cm) would still be consistent with the noble gas data. However, these radii are incompatible with short-lived radionuclide data, which suggest a smaller radius. As discussed later, both data sets can be brought into agreement, if the Stubenberg meteoroid experienced a recent collision which decreased its radius. In the Leya and Masarik model, we hence average the cosmogenic <sup>3</sup>He, <sup>21</sup>Ne, and <sup>38</sup>Ar production rates for all "allowed" sample depths for meteoroids with 35 to 50 cm radius, i.e. where the predicted <sup>22</sup>Ne/<sup>21</sup>Ne matches the measured one within uncertainty. The most likely shielding depth would then be 10–12 cm for the samples analyzed here (Leya and Masarik, 2009). The "uncertainty" of the resulting production rates given in Table 4 covers the maximum and minimum values obtained for all allowed conditions. Note that systematic uncertainties of the Leya and Masarik (2009) production rate model, assumed to be ~15-20 % are not included in the uncertainties given in Table 4.

Data points of both samples plot close to or above the "Bern line" in <sup>3</sup>He/<sup>21</sup>Ne vs. <sup>22</sup>Ne/<sup>21</sup>Ne space (see, e.g., Fig. 5 of Scherer et al., 1998) suggesting that Stubenberg did not lose cosmogenic <sup>3</sup>He during its travel to Earth.

## Radionuclides

The activities for the very short-lived radionuclides are given in Table 5 and the naturally occurring radionuclides (Table S3) are in agreement with the average concentrations in ordinary LL chondrites (Wasson and Kallemeyn, 1988).

#### Oxygen isotopes

The oxygen isotope composition of the analyzed fragment of Stubenberg is  $\delta^{17}O = 3.854 \%$ and  $\delta^{18}O = 5.067 \%$  (Figs. S3 and 8). This gives an offset in  $\delta^{17}O$  of  $\Delta^{'17}O = 1.216 \%$ relative to a reference line with a slope of 0.5305 that passes through San Carlos olivine  $(\Delta^{'17}O = 0 \%)$ . The data for Stubenberg are consistent with those of other LL chondrites.

## Titanium isotopes

The Ti isotope compositions of Stubenberg measured in Münster (n=12;  $\epsilon^{46}$ Ti = -0.08 ± 0.11,  $\epsilon^{48}$ Ti = -0.05 ± 0.11,  $\epsilon^{50}$ Ti -0.67 ± 0.09; all uncertainties are 95% c.i.) and Zurich (n=7;  $\epsilon^{46}$ Ti = -0.10 ± 0.10,  $\epsilon^{48}$ Ti = 0.00 ± 0.03,  $\epsilon^{50}$ Ti -0.81 ± 0.09) are within uncertainty identical to each other, and agree within error with literature values for ordinary chondrites (Trinquier et al., 2009; Zhang et al., 2012; Williams, 2015) (Fig. S4).

#### Chromium isotopes

The Cr isotope composition of Stubenberg is  $\varepsilon^{53}$ Cr = 0.428 ±0.027 and  $\varepsilon^{54}$ Cr = -0.167 ±0.063. The composition of  $\varepsilon^{53}$ Cr is ~8 ppm higher than the previously highest value for ordinary chondrite (OC), while the  $\varepsilon^{54}$ Cr value is within the range of previously reported OC compositions (e.g., Jenniskens et al., 2014; see Fig. 8). Although the Cr isotope composition of Stubenberg is on the extreme of the OC data array its composition falls within the OC region when compared with other meteorite groups.

#### Solvent-soluble organic matter

The analysis of the methanol-soluble organic fraction of Stubenberg by ultrahigh resolution mass spectrometry was undertaken in less than 24 hours after the fragment was found on the field and in less than one week after its fall. For details on the significance of the analysis see Ruf et al. (2017). Care was taken with respect to differentiate indigenous organic matter and possible terrestrial contamination by analyzing corresponding solvent methanolic blanks (Schmitt-Kopplin et al., 2010). The main characteristics of the compositional space was a high density of m/z signals (higher than 60 at each nominal mass) over a mass range from 150 to 650 amu (Fig. 9). These enormous number of m/z signals were annotated to molecular formulas with respect to C-, H-, N-, O-, S- and Mg-bearing compositions. Majorly the CHO, CHOMg and CHNO compositions rule the detected soluble organic space of the Stubenberg meteorite.

### **Physical Characteristics**

### FTIR-Spectroscopy

The analyzed polished thin sections and rough, unprepared surfaces of the Stubenberg samples show similar spectra compared with those obtained from other metamorphosed LL-group chondrites. The FTIR spectra in the mid-infrared of polished thin sections and rough surfaces show very similar features (Figs. 10 and S5). The characteristic Christiansen Feature (CF), a reflectance minimum used to estimate bulk compositions in remote sensing (Salisbury, 1993) is at 7.9-8.1  $\mu$ m. The strongest Reststrahlen Band (RB) is a 'twin-peak' feature at 10.8  $\mu$ m and 11.3  $\mu$ m of two bands with comparable intensity. Further strong RB are overlapping olivine, pyroxene, and plagioclase features found at 9.1, 9.6, 10.3, and 12.0

 $\mu$ m, and at 13.8, 14.6, 15.6, and 16.7  $\mu$ m at longer wavelengths (Baldridge et al., 2009; Hamilton 2000; 2010).

At shorter wavelengths, bands at 2.9  $\mu$ m indicate probably adsorbed terrestrial water. Weak features at 3.4 and 3.5  $\mu$ m could be aliphatic C-H stretching of CH<sub>2</sub> and CH<sub>3</sub>. Bands at 5.7-6  $\mu$ m could be also due to water, but overlap with C=O stretching modes (Kebukawa et al., 2011).

### Magnetic susceptibility

The average values for the magnetic susceptibility (MS) of the samples, corrected for mass and dimensions, are given in Table 6. The correction procedure produces consistent values over 3 orders of magnitude of the sample mass as shown in Fig. S6. For the conversion of volumetric MS instrument readings into MS the bulk density of  $3.29 \text{ g/cm}^3$  was used.

The overall average of MS is about  $4.800 \cdot 10^{-9} \text{m}^3/\text{kg}$  corresponding to  $\log(\text{MS}) = 3.68$  which places Stubenberg at the low end of the range for LL chondrites (Table 6). The 95% confidence interval of  $\pm 0.03$  for  $\log(\text{MS})$  containing both the experimental scattering and the material inhomogeneity is remarkably narrow. The variability of five measurements at different portions of the main mass (M5) is approximately the same as that of those across all samples indicating a quite homogeneous distribution of metal in the breccia.

### DISCUSSION

## Mineralogical properties and classification of Stubenberg

Stubenberg is a breccia based on presented results. The textures of individual fragments are highly recrystallized (Fig. 3) and only a few relic chondrules are visible in thin section (Fig. 3). These observations already point to a high metamorphic grade of the fragments within the rock. The large grain size of plagioclase and the homogeneous compositions of olivines and pyroxenes also support this. These facts clearly indicate that the fragments within Stubenberg are of type 6. Based on the mean olivine and low-Ca pyroxene compositions of Fa<sub>31.4</sub> and Fs<sub>25.4</sub>, respectively, the rock has to be classified as an LL-group ordinary chondrite breccia. Since no fragments of other petrologic type have been observed so far a classification as a LL6 breccia is appropriate. However, Stubenberg does not belong to the complicated breccias like Almahata Sitta (e.g., Bischoff et al., 2010; Horstmann et al., 2010; Goodrich et al., 2014; Horstmann and Bischoff, 2014) or to the complex regolith breccias among the L- and LL-group ordinary chondrites (Bischoff et al., 1993, 2013a, 2006; Metzler et al., 2010, 2011) and

is also different to the LL5-6 breccia Chelyabinsk (Bischoff et al., 2013b; Righter et al., 2015; Morlok et al., 2017). This is consistent with the lack of solar wind implanted noble gases (Fig. 7); thus, the breccia does not contain fragmental debris that was exposed at the surface and hence has to be characterized as a fragmental breccia. This result is also consistent with the observation that no other lithologies beside type 6 fragments were encountered. Considering the onion-shell model for the setting of the parent body no clasts of lower petrologic type (3-5) originated from the upper areas of the parent body were mixed into the breccia during impact-induced lithification (Kieffer, 1975; Bischoff et al., 1983).

Based on the undulatory extinction in olivine and plagioclase and the presence of planar fractures in olivine, the chondrite is weakly shocked (S3). Thus, Stubenberg is an LL6, S3 ordinary chondrite breccia.

Applying the two-pyroxene thermometer implemented in the QUILF 95 program (Andersen et al., 1993) using the compositions listed in Table 2 yields an equilibration temperature of  $914\pm44$  °C. This is consistent with typical values obtained from LL6 chondrites and interpreted as peak metamorphic temperatures (Slater-Reynolds and McSween, 2005). Nipoor pentlandite has been described from several equilibrated LL chondrites and the coexistence of pentlandite with troilite, kamacite, taenite, and potentially tetrataenite indicates disequilibrium among these opaque phases (Michel-Lévy and Bourot-Denise, 1992; Jamsja and Ruzicka, 2010). The low Ni content places the pentlandite outside of the typical compositional range observed in exsolved pentlandite-pyrrhotite assemblages (e.g., Jamsja and Ruzicka, 2010), but thermodynamic modelling suggests that (partial) equilibration at temperatures below 360 °C will produce Fe-rich pentlandite (Waldner and Pelton, 2004). Disregarding the Co content, the pentlandite may have last equilibrated with the troilite at a temperature of about 250 °C or below. Possibly, the sulfide assemblage was produced and incorporated during the brecciation event and internally equilibrated during cooling without significant chemical exchange with the other opaque phases of the rock.

Two different TiO<sub>2</sub> phases (anatase and rutile) could be identified as single grains by their different Raman spectra (Fig. S2). Ramdohr (1973) proposed the following reaction concerning a likely Ti-oxide formation process in ordinary chondrites: Reduction of ilmenite in the presence of carbon phases (carbon, graphite or hydrocarbons), which have been also detected in Stubenberg (see above) to native iron and Ti-oxide according to the reaction:  $2\text{FeTiO}_3 + \text{C} = 2\text{Fe} + 2\text{TiO}_2 + \text{CO}_2$ .

#### **Chemical properties of Stubenberg**

The bulk composition of the Stubenberg meteorite determined so far is very close to the published concentrations of other metamorphosed LL-group ordinary chondrites like Chelyabinsk (Table 3; Fig. 6; Righter et al., 2015). It can be concluded that no anomalies in elemental composition of the Stubenberg meteorite were detected compared with other LL-chondrites. The O-, Ti-, and Cr-isotope compositions of Stubenberg are also typical of LL ordinary chondrites (Figs. 8, S3, and S4).

Exact molecular formula assignments and resulting compositional space of organic constituents show a consistent coverage of ~4600 elementary compositions. In van Krevelentype representations, the chemical space is plotted as H/C vs O/C ratio or m/z ratio (Fig. 9) keeping abundances of the signals in bubble sizes. This allows insights on the chemical properties on the plotted chemical compositions. Information regarding the degree of unsaturation and oxygenation about the respective chemical compositions are observed, detecting a broad range of oxygenation of mainly aliphatic compounds for the Stubenberg soluble organic matter. Compositions with higher degree of oxidation and higher number of nitrogen and sulfur could be observed, as compared to lower shock-metamorphosed ordinary chondrites (Fig. 9; Schmitt-Kopplin et al., 2012; Popova et al., 2013; Jenniskens et al., 2014). High shock-metamorphosed ordinary chondrites show higher nitrogenated chemical compounds (Popova et al., 2013; Jenniskens et al., 2014). In addition, Stubenberg showed an unusual organomagnesium space (CHOMg) with an extremly dense and complete signature with more than 1000 CHOMg compositions, as many as CHO compositions. This reflects a high metamorphic temperature and a complex shock history of the parent body as explained in detail elsewhere (Ruf et al., 2017) as these thermostable metalorganic compounds could be related to the degree of thermal process of a meteorite. From the rapid results obtained on the soluble organic compositions, we conclude that Stubenberg shows characteristics consistent with the complex shock, brecciation, and lithification history of the breccia.

As expected for a freshly fallen LL6 chondrite, the noble gases do not carry significant terrestrial contamination. All precisely measurable heavier noble gas element and isotopic ratios indicate pure meteoritic rather than terrestrial noble gases. The trapped <sup>36</sup>Ar, <sup>84</sup>Kr, and <sup>132</sup>Xe concentrations are extremely low, consistent with Stubenberg being classified in this

work as LL6. For example, element ratios and isotopic compositions (particularly  ${}^{40}$ Ar/ ${}^{36}$ Ar and  ${}^{129}$ Xe/ ${}^{132}$ Xe) are similar to those found in LL5 Olivenza and LL6 Saint Séverin (Alaerts et al., 1979). The trapped gases that remained after thermal metamorphism are similar to phase Q gases (Busemann et al., 2000). The presence of  ${}^{129}$ Xe derived from short-lived  ${}^{129}$ I with a half-life of ~16 Ma ( ${}^{129}$ Xe/ ${}^{132}$ Xe =1.90 and 1.95) indicates closure for Xe loss within the first ~100 Ma.

We obtain a most likely  $4\pi$  CRE age for Stubenberg of  $36\pm3$  Ma assuming a radius of 35-50 cm (covering all shielding conditions "allowed" by the cosmogenic <sup>22</sup>Ne/<sup>21</sup>Ne of 1.0826 determined for Stubenberg; Leya and Masarik, 2009). This age is an average of all six CRE ages obtained for both samples and the three light cosmogenic isotopes <sup>3</sup>He, <sup>21</sup>Ne, and <sup>38</sup>Ar (Table 4). This age is typical for a large fraction of LL chondrites with CRE ages around 30 Ma (e.g., Scherer et al., 1998; Fig. 4 in Herzog and Caffee, 2014). Much larger pre-atmospheric radii, while still possible using noble gas systematics alone, might not be likely, based on the pre-atmospheric mass of 600 kg as estimated with fireball observations (Spurný et al., 2016).

The remarkable agreement between all CRE ages (Table 4) suggests no preferred He loss due to heating in space (as a possible result of the meteoroid's passage close to the Sun), or during atmospheric entry, which could cause preferential He loss. This observation allows us to also determine gas retention ages for Stubenberg, unaffected by late reset, after ejection from the LL parent body. Using the K, U and Th concentrations determined in this work (Table 3), we obtain nominal (maximum) U/Th-<sup>4</sup>He ages of ~3.0 Ga and K-<sup>40</sup>Ar ages of ~4.1 Ga (Table 4). The measured U concentration of 6.3 ppm appears somewhat low (Table 3). If we use a more typical U concentration of 13 ppm (e.g., Wasson and Kallemeyn, 1988), the U/Th-<sup>4</sup>He ages would be around 2.2 Ga.

Such gas-retention ages can be determined under the assumption that significant partial loss after the calculated closure to gas loss did not occur. Such nominal ages have been observed for LL chondrites before, and show generally that the material of Stubenberg has not been shocked severely late in the LL chondrite parent body's history.

<sup>26</sup>Al (Leya and Masarik, 2009), <sup>60</sup>Co (Eberhardt et al., 1963; Spergel et al., 1986), <sup>54</sup>Mn (Kohman and Bender, 1967), and <sup>22</sup>Na (Bhandari et al., 1993) give as possible size range of the parent body (in the sequence of the given isotopes) radii of 20-120 cm, <20 cm, 10-126 cm, and 15-100 cm. Combining all results of these radionuclides we infer a roughly spherical

meteoroid with <20 cm radius, mainly due to the very small value of  ${}^{60}$ Co. This estimate is further confirmed by the fact that the values for both specimens, so different in size, are similar (see Bhandari et al., 1993). The  ${}^{22}$ Na/ ${}^{26}$ Al of (1.5 +- 0.2) is close to the average value for LL chondrites (Bhandari et al., 2002).

The fall of the Stubenberg LL6 chondrite occurred during the current solar cycle 24 maximum, as indicated by the neutron monitor data (Bartol, 2016). The cosmic ray flux was low in the six months before the fall. As a result, the activities for the very short-lived radionuclides are expected to be low (see Table 5), similar to the earlier reported data for Jesenice (Bischoff et al., 2011 and references cited therein). The naturally occurring radionuclides (Table S3) are in agreement with the average concentrations in ordinary LL chondrites (Wasson and Kallemeyn, 1988).

The calculated difference in radius resulting from short-lived radionuclides (<20 cm) and noble gases (>35 cm) could be explained if Stubenberg experienced a complex exposure history, in which most of the cosmogenic noble gases were produced in a >35 cm object in space, before a collision reduced the meteoroid size to <20 cm. Since the <sup>26</sup>Al data are consistent with a larger (20-120 cm radius; Leya and Masarik, 2009) object, the change in radius would have to have happened within no more than a few half-lives of <sup>26</sup>Al ( $t_{1/2} = 717$ ) ka). A complex exposure history for Stubenberg would also seem likely because its CRE age is a factor 3-4 longer than the typical time such a small meteoroid is likely to escape collisional disruption in the asteroid belt, of 9 and 12 Ma for an R = 20 cm and R = 35 cm stony meteoroid, respectively (Farinella et al., 1998). However, there is an additional discrepancy between the size of Stubenberg as implied by short-lived radionuclides (<20 cm) and the estimated mass of the meteoroid implied from fireball observations (ca. 600 kg, R ~35 cm; Spurný et al., 2016). This could potentially be resolved if the Stubenberg meteoroid had a significant macroporosity: as the cosmic-ray-induced production of cosmogenic noble gases and radionuclides depends on the product of radius and density, a lower bulk density would result in a larger physical radius and thus a much larger mass (since M  $\sim R^3$ ; see Welten et al., 2010). At a fragment density of 3.29 g/cm<sup>3</sup> and a macroporosity of 50%, a meteoroid with a size consistent with the measured  $^{60}$ Co activity would have a mass of ~530 kg, consistent with the mass from the fireball estimate. More detailed modeling of the entry fireball, which is beyond the scope of this paper, will be required to resolve this discussion.

## **Physical properties of Stubenberg**

The analyzed polished thin sections and rough, unprepared surfaces of the Stubenberg samples show similar spectra compared with those obtained from other metamorphosed LL-group chondrites (Figs. 10 and S5), Tuxtuac (LL5) and Paragould (LL6). These spectra were acquired by investigators Carle Pieters and Jitendra N. Goswami with the NASA RELAB facility at Brown University. They show essentially similar RB features at 9.5-9.6, 10.3, 10.8 and 11.2-11.3  $\mu$ m as Stubenberg, while the CF is clearly shifted with 8.6  $\mu$ m compared to 8.0-8.1  $\mu$ m for Stubenberg, indicating a lower content in crystallized mesostasis feldspar (Fig. 10) (Salisbury, 1993).

The comparison of the FTIR data with that of heavily shocked LL6 chondrite Chelyabinsk (Morlok et al., 2017) shows a high similarity in band positions and band shapes with the moderately-shocked lithology (Fig. 10). In addition to the four strong bands between 9.6 and 11.3  $\mu$ m, which are also visible in the moderately-shocked lithology of Chelyabinsk, Stubenberg also has a clear feature at 9.1  $\mu$ m, which is only visible as a shoulder in the Chelyabinsk samples. Shock-darkened and impact melt lithologies show a decrease in feature intensities and loss of bands. This implies a low degree of impact shock of the bulk Stubenberg material. Features visible in the 5.7-6.0  $\mu$ m range occurring in the Chelyabinsk sample are only very weak in Stubenberg. This is most probably due to the differences in sample preparation, where the powdered Chelyabinsk material provided higher surface area resulting in stronger minor features compared to the surfaces analyzed of Stubenberg in this work.

The measured magnetic susceptibility data support the classification of Stubenberg as an LL chondrite. Comparing the average values of LL group members given in Rochette et al. (2003) and Macke (2010) the Stubenberg value is significantly lower than the data of LL3 to LL6 meteorites. On the other hand, regarding the limits of variation in the different groups, Stubenberg might fit into LL4 to LL6, with a tendency towards the very special LL7 group (however, represented only by one measured fall, Uden) with a still lower susceptibility value (3.33).

## CONCLUSION

Stubenberg fell on March 6, 2016 and the first pieces were recovered six days later after an exact calculation of the impact area based on instrumental observations taken by the six cameras of the Czech part of the European Fireball Network. Altogether six individuals

weighing in total 1473 g were recovered. It is classified as an LL6, S3 ordinary chondrite fragmental breccia. The bulk composition of the Stubenberg meteorite is very close to element concentrations of other metamorphosed LL-group ordinary chondrites. This is also consistent with O-, Ti-, and Cr-isotope data and IR spectroscopy.

The measured activities of short-lived cosmogenic radionuclides clearly indicate that the analyzed pieces result from a very recent meteorite fall consistent with the Stubenberg fall. A cosmic ray exposure (CRE) age of  $36\pm3$  Ma has been obtained for the meteoroid. Using the K, U, and Th concentrations determined in this work nominal (maximum) U/Th-<sup>4</sup>He ages of ~3.0 Ga and K-<sup>40</sup>Ar ages of ~4.1 Ga are determined.

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**Table 1**: The Stubenberg meteorite finds (GPS coordinates of the locations referring to the WGS84 datum). The polished sections studied in Münster and Jena were prepared from the 1.298 g piece of M1d and M6, respectively. The bulk chemical studies (Brest) and isotopic studies (Zurich) were performed on samples of M1h.

No.	Mass (g)	Date of find	Latitude (N)	Characteristics of the meteorite piece;
110.	1v1a55 (g)	(yyyy/mm/dd)	Longitude (E)	Finder of the specimen
M1	47.88	2016/03/12	48° 17' 42.9" 13° 06' 58.9"	Meteorite was scattered into 17 pieces at the impact (M1a-M1n); Michael Krippner: 6.21 g (M1a), Sabine Gumpenberger: 23.58 g (M1b), Mathias Stricker: 5.42 g (M1e), and seven other finders
M2	7.66	2016/03/23	48° 17' 44.9" 13° 07' 57.7"	Individual meteorite, broken into two (fitting) parts of 7.21 g (M2a) and 0.45 g (M2b) at the impact;Ralph Sporn + Martin Neuhofer
M3	19.24	2016/03/23	48° 17' 43.9" 13° 07' 45.8"	Meteorite fragment with primary and secondary fusion crust; Ralph Sporn + Martin Neuhofer
M4	42.43	2016/03/26	48° 16' 58.5" 13° 07' 25.0"	Complete individual meteorite with signs of flight orientation; Moritz Karl
M5	1320	2016/04/01	48° 18' 20.9" 13° 05' 36.5"	Complete, flight oriented individual meteorite with deep regmaglypts; Ralph Sporn + Martin Neuhofer
M6	35.89	2016/04/03	48° 17' 53.6" 13° 05' 59.9"	Meteorite fragment with fusion crust and broken faces; Dennis Harries
Total	1473		48° 17.7' 13° 07.0'	

wt%	Olivine	Pyroxene	<b>Ca-Pyroxene</b>	Plagioclase	Apatite	Merrillite	Chromite
	n = 54	n = 43	n = 6	n = 42	n = 16	n = 4	n = 14
SiO <sub>2</sub>	36.8	54.3	53.4	65.3	0.19	0.18	< 0.03
TiO <sub>2</sub>	< 0.02	0.20	0.46	0.05	< 0.02	< 0.01	3.4
Al <sub>2</sub> O <sub>3</sub>	< 0.01	0.16	0.47	20.9	< 0.01	< 0.01	5.2
Cr <sub>2</sub> O <sub>3</sub>	< 0.02	0.15	0.81	< 0.01	< 0.01	< 0.01	54.5
FeO	27.8	16.5	6.4	0.53	0.59	0.62	32.7
MnO	0.47	0.46	0.22	< 0.01	< 0.02	n.d.	0.64
MgO	34.0	26.4	16.0	< 0.03	0.15	3.4	1.51
CaO	< 0.01	1.01	21.3	2.20	52.3	46.1	n.d.
Na <sub>2</sub> O	< 0.01	< 0.03	0.52	9.1	0.33	2.43	< 0.01
K <sub>2</sub> O	< 0.01	< 0.01	n.d.	0.92	< 0.01	0.08	< 0.01
P2O5	< 0.04	< 0.01	< 0.01	< 0.02	39.6	46.4	< 0.01
Cl	< 0.01	< 0.01	n.d.	n.d.	5.7	0.48	< 0.01
Total	99.1	99.2	99.6	99.0	98.9	99.7	98.0
Fa	31.4±0.3						
Fs		$25.4 \pm 0.3$	10.3±0.4				
Wo		2.0±0.3	43.9±0.4				
An				$11.1\pm0.4$			
Or				$5.5 \pm 1.2$			

**Table 2**: Chemical composition of main phases in the Stubenberg fragmental breccia. All data in wt% and obtained by electron microprobe. n.d. = not detected; n = number of analyses.

wt%	<b>ICP-AES</b>	<b>ICP-SFMS</b>	ppm	ICP-SFMS
TiO <sub>2</sub>	0.13	0.12	Zr	5.12
Al <sub>2</sub> O3	2.17		Nb	0.419
Fe <sub>2</sub> O3	29.8		Cs	0.0389
MnO	0.36	0.34	Ba	3.6
MgO	24.9		La	0.451
CaO	1.82	1.90	Ce	1.17
Na <sub>2</sub> O	0.98		Pr	0.175
K <sub>2</sub> O	0.1	0.11	Nd	0.858
P2O5	0.31	0.30	Sm	0.272
ppm			Eu	0.0825
Cr	4573		Gd	0.387
Li		2.29	Tb	0.0693
Be		0.0287	Dy	0.46
K		950	Но	0.101
Sc	8.7	9.33	Er	0.294
V	89	80.6	Tm	0.0437
Mn		2606	Yb	0.274
Со	578	536	Lu	0.0405
Ni	12262		Hf	0.144
Cu		73.1	Та	0.0193
Zn		43.54	W	0.121
Ga		5.06	Pb	0.102
Rb		2.88	Th	0.0476
Sr	12	10.88	U	0.0063
Y	2.8	2.97		

**Table 3**: The bulk composition of Stubenberg. Oxides in wt%, allother elements in ppm.

Table 4a. Noble gas	concentrations and	isotope ratios	in Stubenberg: H	Ie, Ne, and Ar.

	mass / mg	<sup>4</sup> He	<sup>3</sup> He/ <sup>4</sup> He	<sup>20</sup> Ne	<sup>20</sup> Ne/ <sup>22</sup> Ne	<sup>21</sup> Ne/ <sup>22</sup> Ne	<sup>36</sup> Ar	<sup>38</sup> Ar/ <sup>36</sup> Ar	<sup>40</sup> Ar/ <sup>36</sup> Ar
Ι	23.31±0.01	1184.7±1.2	618±6	14.53±0.06	$0.8361 \pm 0.0027$	0.9237±0.0023	$1.46 \pm 0.05$	$1.37 \pm 0.04$	3890±140
Π	71.48±0.01	1201.6±2.7	602±5	14.11±0.10	$0.87 \pm 0.04$	0.95±0.04	$1.36\pm0.05$	$1.19\pm0.06$	4060±160

Table 4b. Noble gas concentrations and isotope ratios in Stubenberg: Kr and Xe.

	<sup>84</sup> Kr	<sup>78</sup> Kr/ <sup>84</sup> Kr	<sup>80</sup> Kr/ <sup>84</sup> Kr	<sup>82</sup> Kr/ <sup>84</sup> Kr <sup>84</sup> Kr = 100	<sup>83</sup> Kr/ <sup>84</sup> Kr	<sup>86</sup> Kr/ <sup>84</sup> Kr	<sup>124</sup> Xe/ <sup>132</sup> Xe	<sup>126</sup> Xe/ <sup>132</sup> Xe
Ι	0.733±0.027	$1.68 \pm 0.07$	15.0±0.5	27.7±1.0	28.0±0.9	28.9±1.2	0.73±0.03	0.78±0.06
II	$0.668 \pm 0.019$	$1.88\pm0.03$	14.71±0.17	28.1±0.5	27.6±0.6	28.8±0.4	0.705±0.025	0.821±0.024

Table 4c. Noble gas concentrations and isotope ratios in Stubenberg: Xe continued.

	<sup>132</sup> Xe	<sup>128</sup> Xe/ <sup>132</sup> Xe	<sup>129</sup> Xe/ <sup>132</sup> Xe	$^{130}$ Xe/ $^{132}$ Xe $^{132}$ Xe = 100	<sup>131</sup> Xe/ <sup>132</sup> Xe	<sup>134</sup> Xe/ <sup>132</sup> Xe	<sup>136</sup> Xe/ <sup>132</sup> Xe
Ι	0.95±0.04	8.9±0.4	190±8	15.7±0.9	80±4	38.5±1.8	32.9±1.5
II	$0.904 \pm 0.024$	9.04±0.19	195±3	16.2±0.3	81.3±1.5	39.2±0.7	32.9±0.6

Table 4d. Radiogenic, trapped, and cosmogenic noble gas concentrations and production rates used.

<sup>4</sup> He <sub>rad</sub> <sup>#</sup>	<sup>40</sup> Ar <sub>rad</sub> <sup>#</sup> <sup>36</sup> Ar <sub>tr</sub> <sup>#</sup>	<sup>3</sup> He <sub>cos</sub> <sup>#</sup>	<sup>21</sup> Necos <sup>#</sup>	$^{38}\mathrm{Ar_{cos}}^{\#}$	P3*	P21*	P38*
I 771±33	5690±280 0.18±0.07	73.2±0.7	$15.8\pm0.4$	$1.966 \pm 0.008$	1.957	0.4227	0.0501
<b>II</b> 793±33	5530±310 0.35±0.06	72.3±0.6	15.5±0.3	$1.562 \pm 0.007$	±0.036	$\pm 0.008$	$\pm 0.0011$

**Table 4e.** "Shielding indicator" cosmogenic <sup>22</sup>Ne/<sup>21</sup>Ne, CRE and nominal gas-retention ages.

	( <sup>22</sup> Ne/ <sup>21</sup> Ne)cos	<b>T</b> 3	T <sub>21</sub>	T38	Taverage	U/Th- <sup>4</sup> He	K- <sup>40</sup> Ar
Ι	1.0826±0.0026	37±1	37±1	39±1	36±3	2.99±0.10	4.09±0.08
Π	$1.05 \pm 0.05$	37±1	37±1	31±1		3.06±0.10	4.05±0.09

He, Ne, Ar concentrations in  $10^{-8}$  cm<sup>3</sup>/g STP Kr, Xe concentrations given in  $10^{-10}$  cm<sup>3</sup>/g STP. CRE ages are in Ma, retention ages in Ga. T<sub>average</sub> is the average of all six values including standard deviation. The larger scatter in the <sup>38</sup>Ar ages might be due to variations in Ca. Uncertainties do not include systematic uncertainties of the production rate determinations assumed to be typically 15-20 % and uncertainties in the absolute gas calibrations (2% for He, Ne, Ar and 5% for Kr, Xe, Heber et al. 2009).

#<sup>4</sup>He<sub>cos</sub> subtracted with  $({}^{4}\text{He}/{}^{3}\text{He})_{cos} = 5.65\pm0.45$  (Wieler, 2002),  ${}^{3}\text{He}_{measured} = {}^{3}\text{He}_{cos}$  and  ${}^{21}\text{Ne}_{measured} = {}^{21}\text{Ne}_{cos}$ .  ${}^{38}\text{Ar}_{cos}$  is obtained by decomposition of the measured  ${}^{36}\text{Ar}/{}^{38}\text{Ar}$  into trapped ( ${}^{36}\text{Ar}/{}^{38}\text{Ar} = 5.32 - 5.34$ , air or Q; Busemann et al., 2000) or cosmogenic ( ${}^{36}\text{Ar}/{}^{38}\text{Ar} = 0.63 - 0.67$ ). The variations are covered in the uncertainty of  ${}^{38}\text{Ar}_{cos}$ . The measured  ${}^{40}\text{Ar}$  is assumed to be entirely radiogenic. At maximum 0.9 and 1.8 %, respectively, of the  ${}^{40}\text{Ar}$  in I and II could be atmospheric, assuming  ${}^{36}\text{Ar}_{tr}$  to be entirely air.

\*Production rates were determined based on the most precise shielding indicator, <sup>22</sup>Ne/<sup>21</sup>Ne (here measured = cosmogenic), with the data tables given by Leya and Masarik (2009). We used the more precise ( $^{22}Ne/^{21}Ne)_{cos} = 1.0826\pm0.0026$  from Stubenberg I. The found mass of 1470 g corresponds to a radius of only 4.7 cm, with a density for LL chondrites of 3.29 g/cm<sup>3</sup> (Wilkison and Robinson, 2000). However, the pre-atmospheric mass has been determined to be 600 kg (Spurný et al., 2016). This limits the maximum radius conservatively to 45 cm (corresponding to 1200 kg mass). Based on ( $^{22}Ne/^{21}Ne)_{cos}$  and the Leya and Masarik (2009) model, the minimum radius is 35 cm. The production rates and the uncertainties given in the table cover all values allowed in this range.

Nuclide	Half-life	Stube	Stubenberg				
		M1b (23.58 g)	M5 (1320 g)				
<sup>48</sup> V	15.9735 d	$18 \pm 2$	n.d.				
<sup>51</sup> Cr	27.704 d	$32 \pm 9$	n.d.				
<sup>7</sup> Be	53.22 d	$74 \pm 10$	$85.4\pm8.8$				
<sup>58</sup> Co	70.83 d	$4.1 \pm 0.7$	$4.7\pm0.6$				
<sup>56</sup> Co	77.236 d	$4.0 \pm 0.7$	$4.5\pm0.6$				
<sup>46</sup> Sc	83.787 d	$7.6\pm0.8$	$6.8\pm0.7$				
<sup>57</sup> Co	271.8 d	$4.9\pm0.9$	$6.0 \pm 1.0$				
<sup>54</sup> Mn	312.3 d	$55.1\pm5.7$	$60.9\pm4.7$				
<sup>22</sup> Na	2.60 y	$86.6\pm9.0$	$87.0\pm6.7$				
<sup>60</sup> Co	5.27 y	$0.9 \pm 0.3$	< 0.5				
<sup>44</sup> Ti	60 y	< 4.8	n.d.				
<sup>26</sup> Al	$7.17 \times 10^5$ y	$60.7\pm4.8$	$57.4 \pm 4.4$				

**Table 5**: Massic activities (corrected to the fall of the meteorite March 6<sup>th</sup>, 2016) of cosmogenic radionuclides (in dpm kg<sup>-1</sup>) in the specimens M1b and M5 of the Stubenberg stone measured by non-destructive gamma-ray spectroscopy. Errors include a 1 $\sigma$  uncertainty of 10% in the detector efficiency calibration (n.d. = not determined).

**Table 6**: Magnetic susceptibility (MS) of representative individuals and fragments of the Stubenberg fall (compare Table 1) and of LL meteorites (falls only) from the literature for comparison: (1) Rochette et al., 2003, (2) Macke, 2010. N = number of analyzed LL-chondrites.

Sample	Mass [g] or N	MS [10 <sup>-9</sup> m <sup>3</sup> /kg]	log MS [10 <sup>-9</sup> m <sup>3</sup> /kg]	Ref.
M1b	23.58	4868	3.687	this study
M1c	1.834	4668	3.669	this study
M1n	2.289	4764	3.678	this study
M2a	7.212	4889	3.689	this study
M3	19.236	5076	3.705	this study
M5	1320	4723	3.674	this study
M6	35.887	4728	3.675	this study
Average		$4817 \pm 172$	$3.683\pm0.015$	this study
LL 3	8		4.37 (4.03 - 4.67)	(1)
LL 3	5		4.38 (4.19 - 4.57)	(2)
LL 4	5		4.18 (3.58 - 4.43)	(1)
	5		4.27 (3.61 - 4.66)	(2)
LL 5	12		4.13 (3.70 - 4.54)	(1)
	6		4.21 (3.83 - 4.65)	(2)
LL 6	19		3.95 (3.56 - 4.27)	(1)
	16		3.93 (3.46 - 4.71)	(2)
LL 7	1 (Uden)		3.33	(1)

### **Figure Captions:**

**Fig. 1:** Calculated strewn field near Stubenberg (Germany) with find locations of the six recovered individuals (M1 - M6). The northern part of the strewn field is dominated by forest, in which the samples M5 and M6 were found. Most other areas represent farmland. The river "Inn" marks the border to Austria in the south. Source of base map: Google Earth.

**Fig. 2:** Images of the six Stubenberg meteorites found between March 12 and April 3, 2016. (a) This broken fragment M1b of 23.58 g is the largest part of 17 pieces, totaling 47.88 g. (b) A chip of 0.45 g broke off from the individual meteorite M2 of 7.66 g at the impact in a field. (c) The meteorite fragment M3 weighing 19.24 g was partially buried in the soft ground. (d) Complete individual meteorite M4 of 42.43 g found on the northern bank of the river Inn. (e) The impact hole of the largest Stubenberg individual meteorite M5 of 1320 g in the forest ground was 14 cm deep. (f) The fragment M6 weighing 35.89 g shows fusion crust on the top, and broken faces at the bottom.

**Fig. 3:** Photomicrographs of typical features of the Stubenberg fragmental breccia. (a) highlyrecrystallized texture within an LL6 fragment. (b) Fragmented appearance of the bulk rock. The boundaries of individual fragments are well-observable. (c) Large plagioclase grains are common as well as thin shock veins shown in (d). (e) Occasionally, relict chondrules like this BO-chondrule are visible. (f) Olivines show several sets of planar fractures. Abbreviations: plag = plagioclase, ol = olivine, py = low-Ca pyroxene, ap = apatite, m = metal, tr = troilite, ca-py = Ca-pyroxene. Images in polarized light (a,b,c,f), crossed Nicols (a,c,f); images in back-scattered electrons (d,e).

**Fig. 4**: Pentlandite in the Stubenberg meteorite. a) SEM-BSE overview showing the elongated pentlandite-troilite aggregate. b) SEM-BSE detail showing the intergrowth of pentlandite (Pn) and troilite (Tro). Arrows mark the location of an extracted FIB section. c) TEM-SAED pattern of pentlandite close to zone axis [1-10]. d) TEM-SAED pattern of pentlandite and troilite showing the orientation relationship outlined in the text.

**Fig. 5:** BSE image of the broken surface of fragment M1h (1.652 g) of the Stubenberg meteorite with troilite (white) and abundant olivine (light grey). The dark grey grains represent plagioclase. Note the well-preserved  $120^{\circ}$  triple junctions between the mineral phases.

**Fig. 6:** Bulk composition of Stubenberg in comparison to the LL chondrite Chelyabinsk. Chelyabinsk data from Righter et al. (2015). For normalization the CI-values of Barrat et al. (2012) are used.

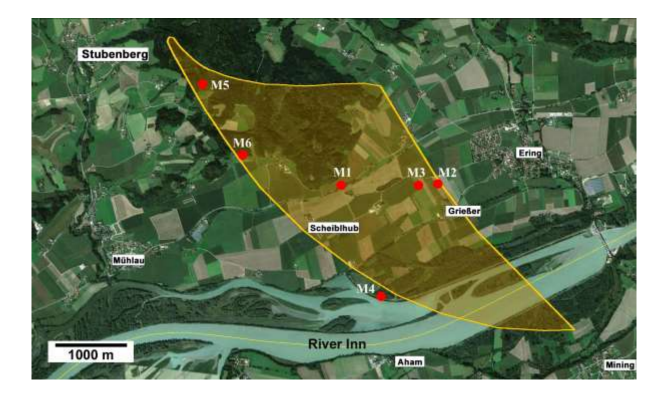
**Fig. 7**: Neon isotopes show that Stubenberg does not contain implanted solar wind noble gases. Both data points plot exactly on the cosmogenic endmember composition, implying that contributions from trapped Ne are negligible. For comparison, data for a selection of LL chondrites that contain mostly trapped Ne (in most cases fractionated solar wind (fSW), are also shown. Typical pure trapped Ne endmembers (air, SW (Heber et al., 2009), and Q (Busemann et al., 2000, 2014) plot in the upper left of the figure). References for LL chondrite literature data: Heymann and Mazor, 1966; Bischoff et al., 1993; Eugster et al., 1993; Scherer et al., 1998; Schultz and Franke, 2004.

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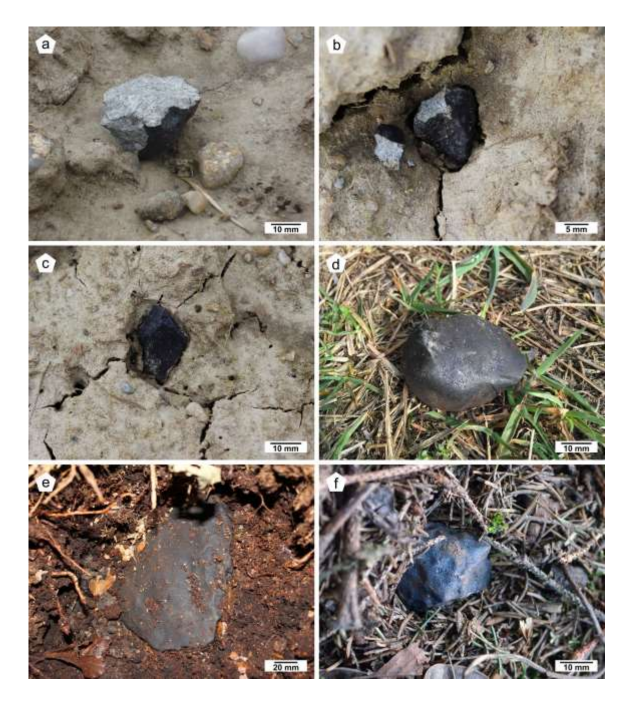
Fig. 9: Negative electrospray ionization mode (ESI) Fourier transform ion cyclotron resonance mass spectrometry ((-) ESI-FT-ICR-MS) FT-ICR-MS analysis of methanolic

extracts revealed ~4600 elementary compositions in the CHNMgOS space. (A) illustrates the mass profile in the range 150 to 650 amu with a zoomed-in detail on the high chemodiversity, as shown exemplary for the nominal mass 319. This enlarged mass range shows more than 70 m/z signals. Panel (B), (C): The number of nitrogen and sulfur atoms in the molecular compositions is shown as a function of the number of carbon atoms. Addititionally, the soluble organic compositions are depicted in van Krevelen diagram-type representations of the data with additional extension of the CHNOS space (D and E) and the CHOMg space (G). The bubble size represents the relative intensity of the mass peaks and each data point is related to one molecular formula. Panel (F) depicts the distribution of the chemical subspaces CHOMg, CHO, CHOS, CHNO and CHNOS.

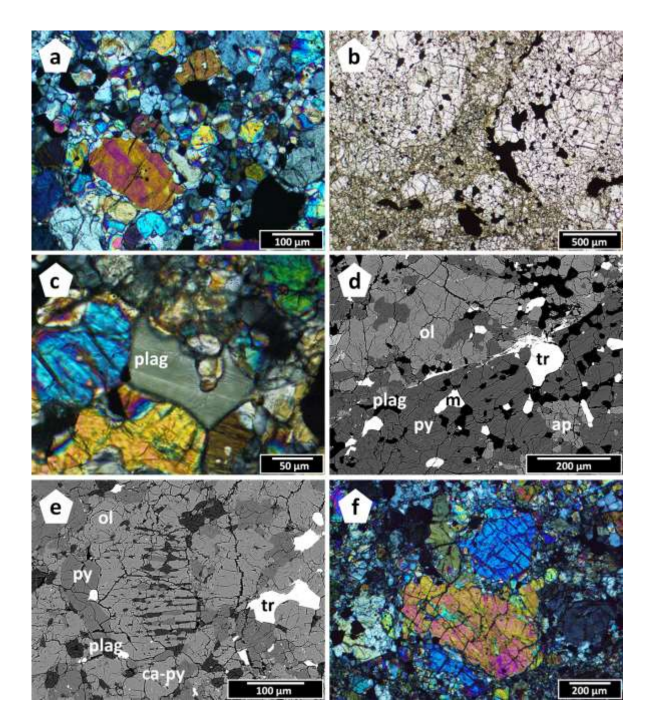
**Fig. 10**: Mid-infrared spectrum of Stubenberg (in black) compared to other LL chondrites Tuxtuac and Paragould (on top) and various Chelyabinsk lithologies (bottom; Morlok et al., 2017). Vertical lines mark characteristic features. Stubenberg does not show the loss of feature strengths exhibited by the Chelyabinsk lithologies resulting from shock metamorphism, thus indicating a generally low degree of impact shock.



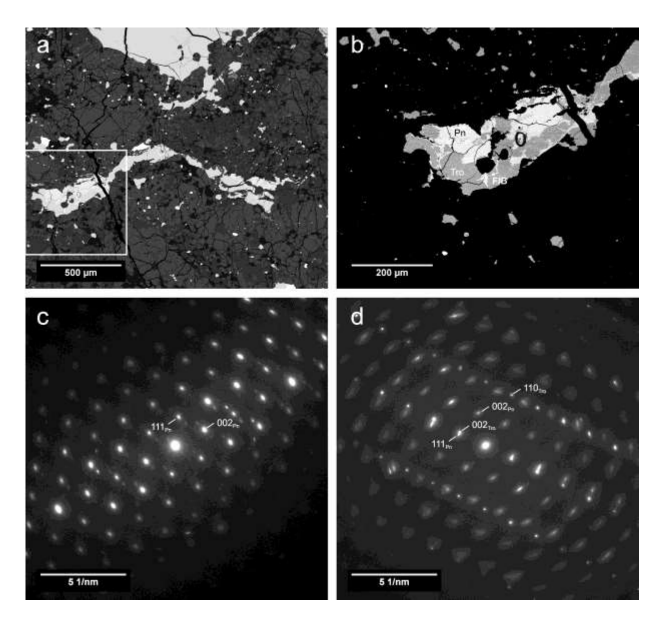
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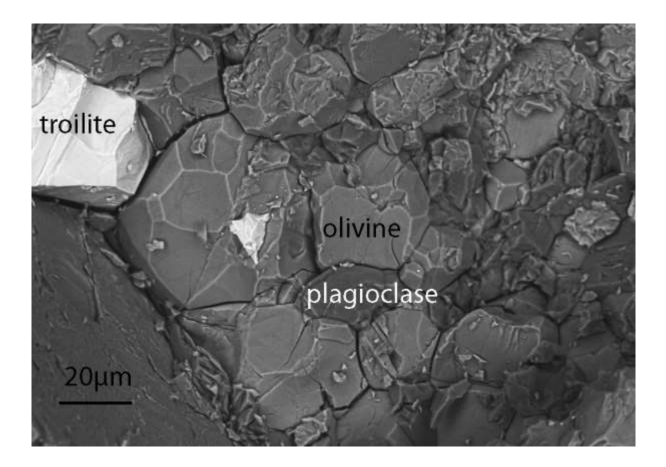
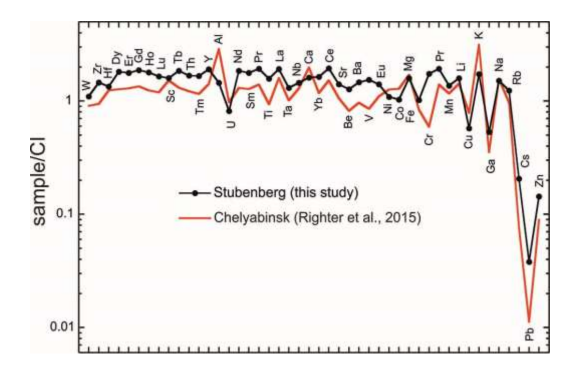
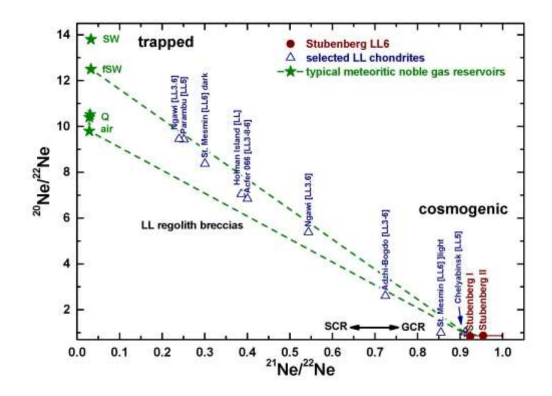


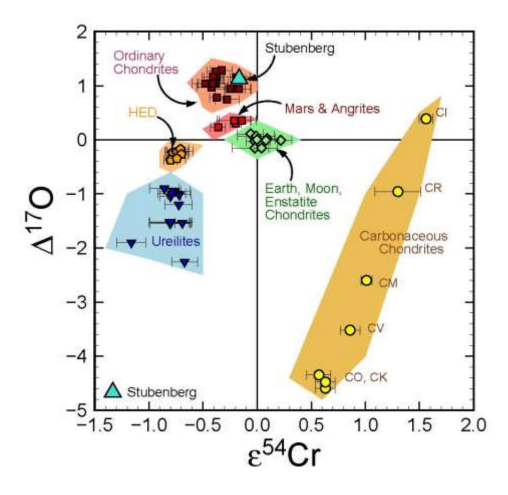
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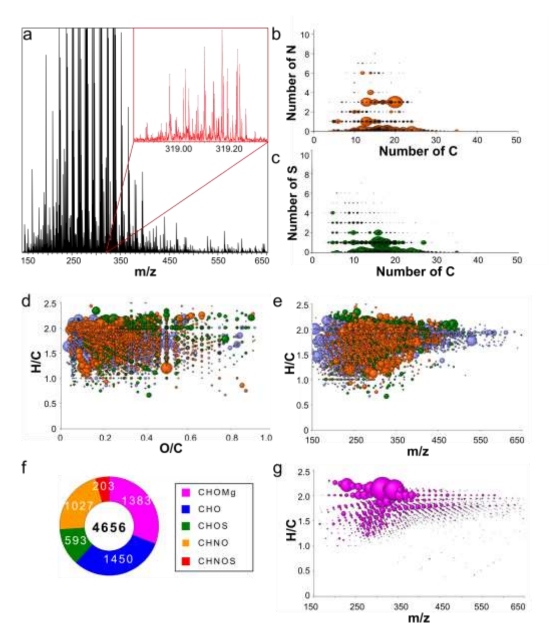
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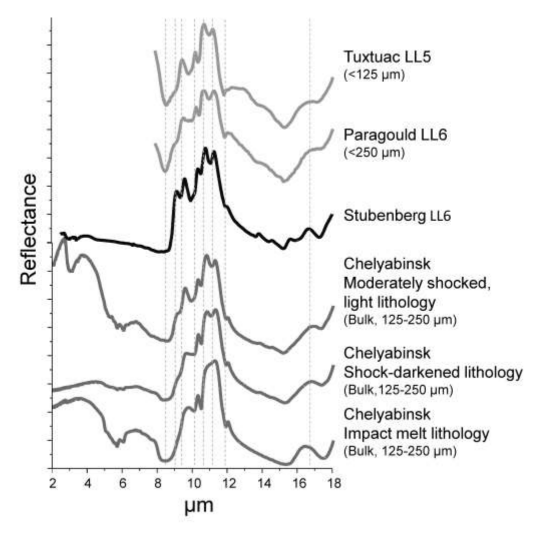
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# **Supplemental Materials**

## Samples and Analytical procedures -

### Optical and electron microscopy

Several thin and thick sections of Stubenberg (PL16031, PL16032, and PL16033 (thin sections University of Münster); H130,4a (thick section) and H130,4b (thin section) (University of Jena)) were studied by optical and electron microscopy. For optical microscopy in transmitted and reflected light a ZEISS polarizing microscope (Axiophot) was used.

A MIRA3 TESCAN field emission electron microscope at the University of Prague was used to resolve the fine-grained mineral assemblages. In Münster a JEOL 6610-LV electron microscope (SEM) at the Interdisciplinary Center for Electron Microscopy and Microanalysis (ICEM) at the Westfälische Wilhelms-Universität Münster was used to study the fine-grained textures of and to identify the different mineral phases. At the University of Jena a FEI Quanta 3D FEG equipped with an EDAX EDS system was used for mineral characterization and determination of modal abundances via BSE imaging and X-ray elemental mapping. A sulfide sample was extracted for transmission electron microscopy (TEM) using focused ion beam (FIB) preparation (Harries and Langenhorst 2013). The FIB sample was studied using a 200 keV FEI Technai G2 equipped with an Oxford SDD EDS detector. In Munich SEM investigations were conducted with a Zeiss EVO MA10 environmental scanning electron microscope, using a high voltage of 15 kV. EDX measurements were performed with an Energy Dispersive X-ray Spectrometer Bruker AXS XFlash 410/M. Fragment M1h (1.652 g) was investigated without any contacting, coating, or sputtering.

## Mineral analyses

Most quantitative mineral analyses were obtained using a JEOL JXA 8900 Superprobe electron microprobe (EPMA) at the ICEM, which was operated at 15 kV and a probe current of 15 nA. Natural and synthetic standards were used for wavelength dispersive spectrometry. Jadeite (Na), kyanite (Al), sanidine (K), chromium oxide (Cr), San Carlos olivine (Mg), hypersthene (Si), diopside (Ca), rhodonite (Mn), rutile (Ti), fayalite (Fe), apatite (P), celestine (S), and nickel oxide (Ni) were used as standards for mineral analyses. Penlandite was analyzed at the University of Jena using a JEOL JXA 8230 Superprobe operated at 20 kV and 15 nA. Standards used were pyrrhotite (S, Fe) and pure metals (Ni, Co). Using the Münster-SEM for quantitative analysis, samples and appropriate mineral standards were measured at an excitation voltage of 20 kV, and the beam current constancy was controlled by a Faraday cup. The attached EDS system was used for chemical characterization and analyses of the different mineral constituents. Standard (Astimex) olivine (Mg, Fe, Si), jadeite (Na), plagioclase (Al), sanidine (K), diopside (Ca), rutile (Ti), chromium-oxide (Cr), rhodonite (Mn), and pentlandite (Ni) were used as natural and synthetic standards. For these EDS analyses the INCA analytical program provided by Oxford Instruments was used.

### Raman Spectroscopy

Raman spectroscopic investigations were conducted on the following fragments:

M1c (1.834 g), M1e (5.418 g), M1h (pieces of 1.652 g fragment), M1m (0.418 g) and M2b (0.45 g). A Horiba Xplora Integrated confocal Laser micro Raman system on an Olympus BX51 microscope was used for our investigations. Magnifications were between 100 and 1000x (Long Distance objectives 10x and 100x) with acquisition times of 3-5 s and accumulation numbers of 2-5. The software used was LabSpec 5.78.24 provided by Horiba. The Raman experiments have been performed exclusively on non-prepared samples because in this way we can guarantee to investigate only pristine material excluding any effects of sample preparation/polishing or sputtering (coating).

High resolution mapping was performed in order to detect and identify all potentially present phases and also exsolution/zonation-effects. Point measurements were conducted on phases of special interest. Most Raman spectra have been obtained using a 532nm Nd/YAG Solid State Laser, in a few cases also the 685 nm Laser was used (for example to suppress higher fluorescence background). Laser power on the sample surface was 8-10 mW, grating was set to 1800. Optical resolution (Laser spot size) was 0.7  $\mu$ m, the Raman spectra precision was +/-1-2 cm<sup>-1</sup>.

The following mapping settings have been used:  $3 \times 3$  up to  $10 \times 10$  maps, point distances of  $1\mu$ m (individual point spectra overlap reaching higher precision). Measurements were conducted from 50 - 2500 cm<sup>-1</sup>, in some cases from 50 - 4000 cm<sup>-1</sup>. Baseline correction was performed using degree 6 or 8 polynomials. Peak search and fitting was done with LabSpec "Peak Search" program. Calibration was done manually using Si calibration standards; additional calibration tests were performed using other mineral standards (e.g., olivines from San Carlos, Iceland calcite or standard graphite).

### Bulk chemical analysis

The sample of about 0.5 g from Stubenberg fragment M1h was crushed and homogenized. This homogenized sample was used for bulk chemical and isotopic studies in Plouzané, Zurich, and Münster. The chemical composition of the bulk sample was obtained by using ICP-AES and ICP-SFMS. The bulk composition using ICP was performed at the Institut Universitaire Européen de la Mer, Université de Bretagne Occidentale in Plouzané, France.

#### Ti isotope analyses

Titanium isotope compositions were measured using Thermo Fisher Scientific Neptune Plus MC-ICPMS at the Institut für Planetologie in Münster and the Institute of Geochemistry and Petrology at ETH Zurich.

For Zurich: A 50 mg sample aliquot in powder form was dissolved in a 6 ml square body Savillex Teflon beakers using HF:HNO<sub>3</sub> (ratio 3:1) in an oven at 160 °C for 3-4 days (after Prytulak et al., 2011). A three-stage ion exchange procedure for Ti purification was adapted from Williams (2015) with a modification based on Zhang et al. (2011). For the first column, the procedure of Zhang et al. (2011) using TODGA resin was used. This was followed by two repeats of an anion exchange column (Bio-Rad AG1-X8) using 4 M HF for sample loading and matrix elution followed by the elution with 0.5 M HCl + 0.5 M HF and the collection of Ti in 6 M HCl + 1M HF (Schönbächler et al., 2004; Williams, 2015). The Ti isotope analyses were performed using a Neptune Plus MC-ICPMS at ETH in high and medium resolving power mode in conjunction with a Cetac Aridus II desolvating system. The samples were bracketed by an in-house Ti wire standard solution (Williams, 2015). Each sample measurement included one block with 40 integrations of 8.39 seconds for the main and 4.19 seconds for the second line, which is necessary to correct for isobaric interferences. On-peak background corrections were applied and the data was internally normalized to <sup>49</sup>Ti/<sup>47</sup>Ti =0.749766 (Niederer et al., 1981). The sample uncertainty is reported as standard error based on repeated analyzes of the sample solution obtained on three different days.

For Münster: About 50 mg of the sample powders were dissolved in 15 ml Savillex Teflon beakers on a hotplate at 170 °C for several days with a mixture of HF:HNO<sub>3</sub> (ratio 3:1) and aqua regia, and Ti was separated from the matrix broadly following previously established procedures (Zhang et al., 2011). Titanium isotope measurements were made using Thermo Scientific Neptune Plus MC-ICPMS in high resolving power mode (Zhang et al., 2011).

Solutions containing about 200 ppb Ti were introduced through a Cetac Aridus II desolvating system, resulting in a  $\sim 4 \times 10^{-10}$  A ion beam on <sup>48</sup>Ti. Measurements consisted of a 30 s baseline measurement (deflected beam) followed by 20 isotope ratio measurements of 8 s each. Mass bias was corrected using the exponential law and <sup>49</sup>Ti/<sup>47</sup>Ti = 0.749766. Titanium isotope anomalies are reported as parts per ten thousand deviation ( $\epsilon$ -notation) from the terrestrial OL-Ti (Millet and Dauphas, 2014) bracketing standard. The sample uncertainty is reported as standard error (student-t 95% confidence interval) based on repeated analyzes of the sample solution obtained on three different days.

### Cr isotope analyses

Chromium isotope compositions were determined by MC-ICPMS at the Institute of Geochemistry and Petrology at ETH Zurich and are described in more detail in Steele and Schönbächler (2016). An aliquot of the same dissolution analyzed for Ti isotopes in Zurich, was also analyzed for Cr isotope composition. It was verified that the dissolution method of Prytulak et al. (2011) dissolves refractory phases such as spinel and chromites, which contain significant amounts of Cr and are not readily dissolved during hotplate dissolution. Chromium was separated from the sample matrix using a three stage ion exchange procedure. A first column, developed from Trinquier et al. (2008), used AG-50W cationic resin and 1 M HCl to isolate Cr from the bulk of the matrix. Two subsequent columns adapted from Steele et al. (2011) were used to separate Cr from direct isobaric interferences (Tru.Spec, Ti, V, and Fe), residual matrix and organic species (AG-50W). Chromium isotope compositions were determined using a Neptune Plus MC-ICPMS in high-resolving power mode (M/ $\Delta$ M > 9000) to resolve isobaric interferences from the plasma gasses (ArC, ArN, and ArO). Samples were introduced as 6 ppm solutions in 0.4 M HNO<sub>3</sub> using a Cetac Aridus II desolvating nebuliser and yielded a  $\sim 1.5$  nA <sup>52</sup>Cr ion beam. Each sample measurement was bracketed by an on peak blank measurement, which was subtracted, and a Cr NIST SRM 979 standard measurement (also blank corrected). A sample or standard measurement consisted of 80 cycles of 16 s integrations and blank measurements were 40 cycles of 8 s integrations. Samples and standards were corrected for natural and instrumental mass-dependent isotope fractionation using the exponential law (Russell et al., 1978) and  ${}^{50}Cr/{}^{52}Cr = 0.051859$  (Shields et al., 1966). The isotope composition is reported as the parts per ten thousands deviation from the Cr NIST SRM 979 standard. The sample was measured 3 times and the isotope composition and uncertainty are the average and standard error of these repeat measurements. The

uncertainty was corrected for small sample bias using the method of Gurland and Tripathy (1971) and is in agreement with the long-term external precision.

### Noble gas studies

The noble gases He-Xe in two bulk samples "I" (23.3 mg) and "II" (71.5 mg) were analyzed at ETH Zurich. We extracted the gases in one temperature step at ~1700 °C, with the standard procedures described in detail by Riebe et al. (2017, in revision). Blank corrections were mostly negligible (<0.3% for all He, Ne, and Ar isotopes), but 7 and 19% for <sup>84</sup>Kr, and 3 and 8 % for <sup>132</sup>Xe for the smaller (I) and larger (II) sample, respectively. A re-extraction step at slightly elevated temperature (1750 °C) showed that sample I was completely degassed in the main step. Concentrations and isotope ratios are given in the Appendix of Table 4.

## Radionuclides

Cosmogenic radionuclide concentrations have been analyzed by means of non-destructive high purity germanium (HPGe) gamma spectroscopy. The counting efficiencies have been calculated using thoroughly tested Monte Carlo codes. The specimen M1b (23.58 g) was measured in the underground laboratories at the Laboratori Nazionali del Gran Sasso (LNGS) (Arpesella, 1996), whereas piece M5 (1320 g) was measured at the low level laboratory of the Max-Planck Institut für Kernphysik (Heusser et al., 2015). The specimen M1b was received at LNGS already 14 days after the fall, so that also rather short-lived radionuclides such as <sup>48</sup>V, and <sup>51</sup>Cr could be measured.

### Oxygen isotope analyses

The oxygen isotope composition of Stubenberg was analyzed by means of laser fluorination in combination with gas source mass spectrometry (Sharp, 1990). The sample was loaded along with San Carlos olivine in a 19-pit stainless steel sample holder. We used a 2-window (quartz glass viewing port and BaF<sub>2</sub> for the CO<sub>2</sub> laser beam) sample chamber as described in Sharp (1990). The samples were fluorinated with ~ 40 mbar BrF<sub>5</sub>. The unwanted gaseous reaction products (mainly BrF<sub>5-x</sub> and SiF<sub>4</sub>) were removed my means of cryogenic trapping. The sample O<sub>2</sub> was collected at -196 °C on 5 Å molecular sieve for 7 min. For further purification (removal of N<sub>2</sub> and NF<sub>3</sub>), sample O<sub>2</sub> was transported with a He carrier gas stream through a 3 m long packed molecular sieve gas chromatography column (Restek) that was kept at 50 °C. The sample O<sub>2</sub> was then collected downstream in a second, low volume molecular sieve trap. After evacuation of He from the second trap, sample O<sub>2</sub> was expanded to the bellows of a Thermo MAT253 gas source mass spectrometer. The sample gas was analyzed relative to a working gas for ~ 60 minutes (90 cycles). The external reproducibility of San Carlos olivine measurements was 0.1 ‰ for  $\delta^{18}$ O and 0.006 ‰ for  $\Delta^{17}$ O (1 $\sigma$  SD, single analysis). The data were normalized to the composition of San Carlos olivine ( $\delta^{17}O_{VSMOW2} = 2.715$ ,  $\delta^{18}O_{VSMOW2} = 5.220$  ‰; see Pack et al., 2016).

### FTIR Spectroscopy

Measurements were made using a Bruker Vertex 70v at the IR/IS laboratory of the Institut für Planetologie in Münster. We analyzed polished thin sections and rough, unprepared surfaces of meteorite sample. All analyses were collected using identical incidence and emergence angles  $(30^{\circ}/30^{\circ})$ , using a Bruker A513 bidirectional reflectance stage. For each individual spectrum, 512 scans were accumulated in the wavelength range from 2 to 18 µm. All reflectance analyses were made at low pressure  $(10^{-3} \text{ bar})$ . For background calibration, either a polished gold mirror (for thin section analyses) or a diffuse reflectance standard (rough surfaces) was used. The spectra will be made available as part of the Berlin Emission Database (BED), a mid-infrared spectral database to provide data for the MERTIS (MErcury Radiometer and Thermal Infra-red Spectrometer) mid-infrared spectrometer on the upcoming ESA/JAXS BepiColombo mission to Mercury and later missions (Benkhoff et al., 2010; Hiesinger et al., 2010).

### **Organics**

A destructive methanolic solvent extraction of organics matter was performed in a pillar using 50 mg freshly broken fragment of the Stubenberg meteorite (from sample M1j). All precautions were taken to avoid external organic contamination and the broken parts were washed intensively with the solvent prior crashing. Negative electrospray ionization mode (ESI) Fourier transform ion cyclotron resonance mass spectrometry (FT-ICR-MS) was undertaken using standard operation procedure as described previously by Schmitt-Kopplin et al. (2010, 2012) and 3,000 scans were averaged. The polar, protic solvent extraction (methanol) and mild ESI-ionization preferentially extracted and ionized polar, oxygenized molecules. Data processing was carried out using the Compass Data Analysis 4.0 (Bruker, Bremen, Germany) and molecular formulas were assigned by a custom made software (NetCalc network), as described by Tziotis et al. (2011). Assigned molecular formulas were

classified into CHO, CHNO, CHOS, CHNOS, and CHOMg molecular series, which were visualized in the class-selective van Krevelen diagrams (Schmitt-Kopplin et al., 2010, 2012; Hertkorn et al., 2015; Ruf et al., 2017).

# **Physical Properties**

Magnetic susceptibility (MS) was determined for seven meteorites and fragments of the Stubenberg fall. Dependent on size and geometry up to five measurements per sample were performed with the SM30 of ZH Instruments.

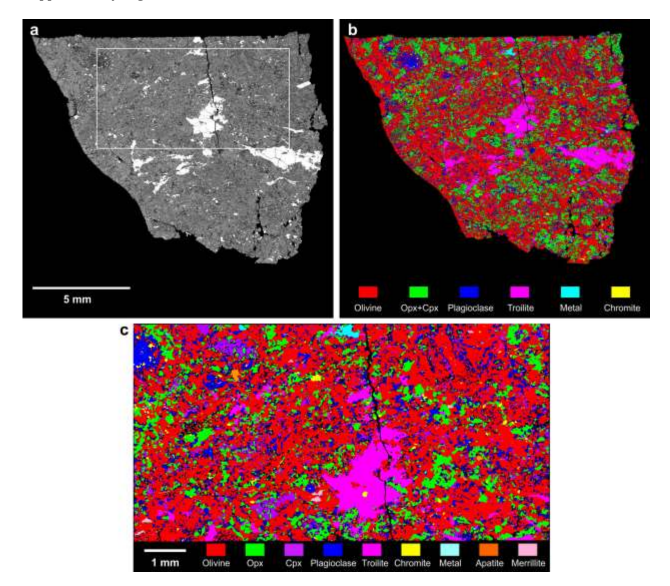
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Supplementary Figures:



**Fig. S1**: Modal analysis of the Stubenberg meteorite. a) SEM-BSE mosaic of sample H130,4a. b) Color-segmented phase map of the SEM-BSE mosaic indicating the different mineral phases. Pyroxenes could not be distinguished based on the BSE coefficients. c) SEM-EDX phase map of the area marked in a. orthopyroxene (Opx) and clinopyroxene (Cpx) are distinguished based on their Ca contents.

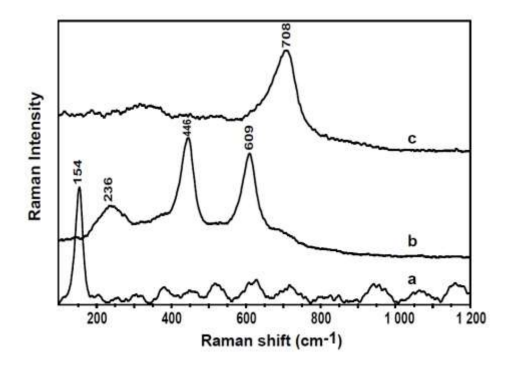
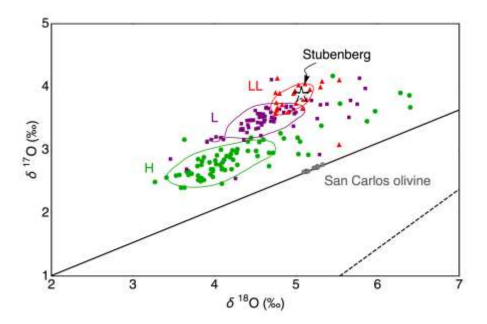
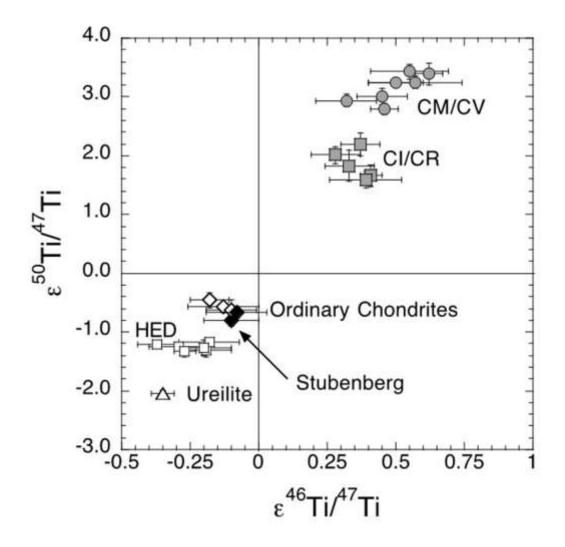


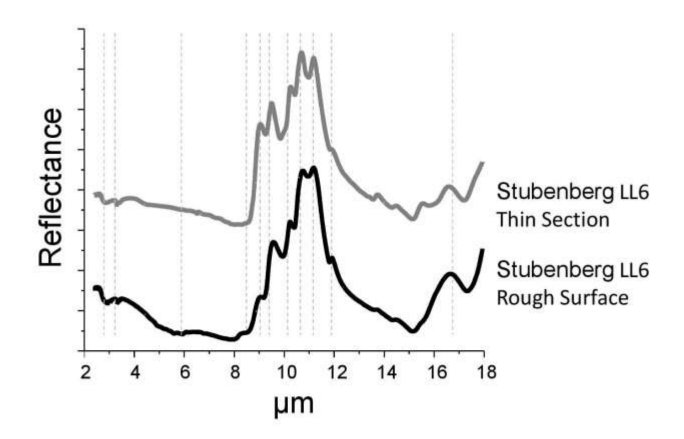
Fig. S2: Raman spectra of anatase (a), rutile (b), and ilmenite (c) from Stubenberg meteorite.



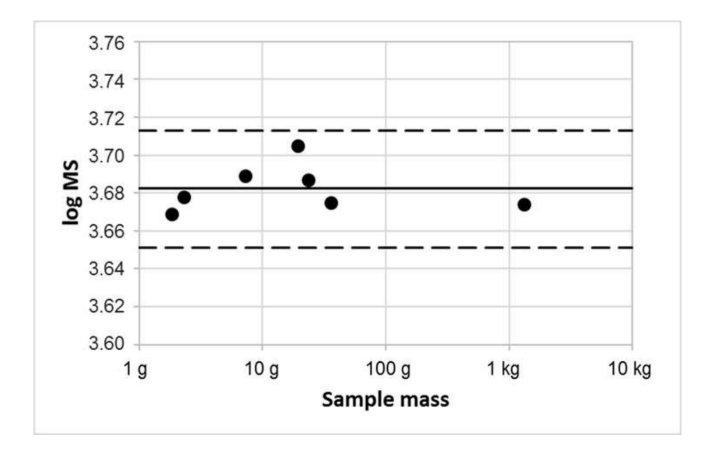
**Fig. S3**: Plot of  $\delta^{17}$ O vs.  $\delta^{18}$ O of Stubenberg (star) in comparison with published oxygen isotope data of H, L, and LL ordinary chondrites. The results of San Carlos olivine measurements (gray filled circles) are displayed along with the terrestrial fractionation line (solid) and the carbonaceous chondrite anhydrous minerals mixing line (dashed).



**Fig. S4**: The Ti isotope compositions ( $\epsilon^{46}$ Ti/<sup>47</sup>Ti vs.  $\epsilon^{50}$ Ti/<sup>47</sup>Ti) for various meteorite groups. Each group occupies a unique space in the diagram. Black diamonds denote Stubenberg. Data from this study and Williams (2015).



**Fig. S5**: Mid-infrared spectra of Stubenberg (in wavelength). Vertical lines mark characteristic features discussed in the text.



**Fig. S6:** Magnetic susceptibility values of representative meteorites and fragments of the Stubenberg fall with average (solid line) and confidence interval (dashed lines at +/-2 standard deviations of all measured values).

Area %	EDS	(normalized)	BSE	(normalized)	
(= <b>vol%</b> )	A =	$A = 50.6 \text{ mm}^2$		$A = 120.5 \text{ mm}^2$	
Olivine	50.71	55.82	45.71	53.03	
Pyroxene			20.48	23.76	
Low-Ca pyroxene	15.14	16.67			
Plagioclase	9.39	10.34	11.59	13.44	
Troilite	8.48	9.34	7.44	8.63	
Diopside	5.71	6.29			
Chromite	0.64	0.70	0.45	0.52	
Metal	0.38	0.42	0.52	0.60	
Merrillite	0.23	0.25			
Apatite	0.16	0.18			
Total	90.84	100.00	86.19	100.00	

**Table S1:** Modal mineral abundances of the Stubenberg breccia (sample M6) obtained from EDS X-ray maps and BSE images. A = area used for image segmentation.

wt%	Troilite	Taenite	Kamacite	Pentlandite
	n = 67	n = 38	<b>n</b> = <b>8</b>	n = 6
Fe	62.0	53.2	87.3	49.4
Ni	< 0.02	44.3	3.9	16.3
Со	n.d	1.78	6.1	0.4
S	35.9	< 0.01	< 0.01	34.2
Р	< 0.01	< 0.01	< 0.02	n.d.
Cr	< 0.02	< 0.02	< 0.03	n.d.
Mn	< 0.02	< 0.01	< 0.01	n.d.
Cu	< 0.03	0.22	n.d.	n.d.
Total	97.9	99.6	97.3	99.3

**Table S2:** Chemical composition of metals and troilite within the Stubenberg fragmental breccia. All data in wt%. n.d. = not detected; n = number of analyses.

**Table S3**: Concentration of primordial radionuclides (ng  $g^{-1}$  for U and Th chains and  $\mu g g^{-1}$  for  $K_{nat}$ ) in the specimens M1b and M5 of the Stubenberg stone measured by non-destructive gamma-ray spectroscopy. Errors include a 1 Sigma uncertainty of 10% in the detector efficiency calibration.

Nuclide	Stubenberg			
	M1b (23.58 g)	M5 (1320 g)		
<sup>238</sup> U	$16 \pm 2$	$13 \pm 1$		
<sup>228</sup> Th	$35 \pm 3$	$44 \pm 3$		
<sup>40</sup> K	$830\pm80$	$750\pm80$		