

# **Osmium Isotope Systematics of Ureilites**

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## Abstract

The  $^{187}\text{Os}/^{188}\text{Os}$  for twenty-two ureilite whole rock samples, including monomict, augite-bearing, and polymict lithologies, were examined in order to constrain the provenance and subsequent magmatic processing of the ureilite parent body (or bodies). The Re/Os ratios of most ureilites show evidence for a recent disturbance, probably related to Re mobility during weathering, and no meaningful chronological information can be extracted from the present data set. The ureilite  $^{187}\text{Os}/^{188}\text{Os}$  ratios span a range from 0.11739 to 0.13018, with an average of  $0.1258 \pm 0.0023$  ( $1\sigma$ ), similar to typical carbonaceous chondrites, and distinct from ordinary or enstatite chondrites. The similar mean of  $^{187}\text{Os}/^{188}\text{Os}$  measured for the ureilites and carbonaceous chondrites suggests that the ureilite parent body probably formed within the same region of the solar nebula as carbonaceous chondrites. From the narrow range of the  $^{187}\text{Os}/^{188}\text{Os}$  distribution in ureilite meteorites it is further concluded that Re was not significantly fractionated from Os during planetary differentiation and was not lost along with the missing ureilitic melt component. The lack of large Re/Os fractionations requires that Re/Os partitioning was controlled by a metal phase, and thus metal had to be stable throughout the interval of magmatic processing on the ureilite parent body.

## 1. Introduction

The ureilites represent the second largest group among the achondritic meteorites, comprising ~16% of all achondrites. As of May 2006, the Meteoritical Bulletin database listed about 200 individual ureilite samples, some of which may be paired. Ureilites are best known as a distinct class of achondrites that appear to be products of a significant degree of planetary igneous differentiation, but also preserved primordial signatures that distinguishes them from other achondrite groups. Unlike other achondrite groups, the ureilites do not fall on a mass-dependent isotopic fractionation trend in oxygen isotope space (Clayton and Mayeda, 1988, 1996), but rather fall along a slope  $\sim 1$  line defined by carbonaceous chondrite anhydrous minerals (CCAM). Ureilites also exhibit  $^{33}\text{S}$  isotope anomalies (Farquhar et al., 2000), as well as large differences in  $\delta^{15}\text{N}$  between ureilitic diamond, graphite, and silicate phases (Rai et al., 2003a). These isotopic heterogeneities are thought to preserve the primordial nebular heterogeneity in the accreting ureilite precursor materials and thus argue against a global magmatic equilibration of the ureilite parent body. Another apparently nebular signature that seems to preclude the significant outgassing that is expected from extensive magmatic processing is the high abundance of fractionated primordial noble gases that reside mostly in diamond (Begemann and Ott, 1983; Göbel et al., 1977; Rai et al., 2003b).

Petrographically, the monomict ureilites consist predominantly of olivine, pyroxene (mostly pigeonite, but also orthopyroxene and augite) and  $\leq 10\%$  dark interstitial material consisting of carbon polymorphs (graphite, diamond, lonsdaleite, chaoite), metal, sulfides, and fine grained silicates (Goodrich, 1992; Goodrich et al., 2004; Mittlefehldt, 2003; Mittlefehldt et al., 1998). This material also occurs as veins that intrude the sili-

cates along fractures and cleavage planes. With carbon abundances up to 6-7 wt%, the ureilites are the most carbon-rich meteorite group (Grady and Wright, 2003). On the basis of their silicate mineralogy, the monomict ureilites are classified into three different types: olivine-pigeonite, olivine-orthopyroxene, and augite-bearing. Succinct descriptions of the classification of monomict ureilites are given in Goodrich et al. (2004, 2006). Most monomict ureilites show coarse-grained igneous textures with mineral grains joining in abundant 120° triple junctions. Equilibration temperatures for ureilites estimated from two-pyroxene thermometry range from ~1200° to 1280°C (Chikami et al., 1995; Sinha et al., 1997; Takeda, 1987; Takeda et al., 1989), whereas olivine-pigeonite-liquid thermometry (Singletary and Grove, 2003) suggests a slightly broader range from ~1150° to 1300°C. Monomict ureilites lack plagioclase and are depleted in incompatible lithophile elements. In these respects the ureilites resemble typical terrestrial ultramafic upper mantle rocks (such as lherzolites and harzburgites) that experienced removal of a basaltic component during partial melting events.

The two main origins that have been considered for ureilites are as partial melting residues (Scott et al., 1993; Warren and Kallemeyn, 1992), or as cumulates (Berkley et al., 1976; Berkley and Jones, 1982; Berkley and Keil, 1980). Although most workers now accept the theory that ureilites are residues rather than cumulates, the augite-bearing ureilites, which represent a small percentage of all ureilites, are likely to be cumulates or paracumulates (Goodrich et al., 2004). The olivine and pyroxene core mg# [molar Mg/(Mg+Fe)] of the monomict ureilites as a whole have been proposed to be genetically linked via pressure-dependent carbon redox control (Berkley and Jones, 1982; Goodrich et al., 1987a; Singletary and Grove, 2003, 2006; Sinha et al., 1997; Walker and Grove,

1993). In this ‘smelting’ model, silicate FeO reacts with carbon in the presence of a silicate melt to form Fe-metal and carbon monoxide. A rapid, CO-gas driven localized melt extraction and loss of the melt into space resulting from high eruption velocities could provide an explanation for the lack of basaltic ureilites in the meteorite collections (Keil and Wilson, 1993; Scott et al., 1993; Warren and Kallemeyn, 1992). One problem of the smelting models is that they imply the loss of a significant amount of metallic iron in the more ‘smelted’ i.e. MgO-rich ureilites (Mittlefehldt et al., 2005; Warren and Huber, 2006). The high abundances and chondritic interelement ratios of several highly siderophile (iron-loving) elements (HSE – Ru, Rh, Pd, Re, Os, Ir, Pt, and Au) (Boynton et al., 1976; Higuchi et al., 1976; Janssens et al., 1987; Mittlefehldt et al., 2005; Wänke et al., 1972; Warren et al., 2006), however, are inconsistent with extensive removal of Fe-metal (as expected from the smelting model, or the segregation of a metallic core) because the metal would also effectively scavenge the siderophile elements from the rock.

In this paper, the Re-Os isotope system is used to address open questions in ureilite research. One question concerns the provenance of the ureilite protolith material. A basic tool for distinguishing between different meteorite classes is a categorization based upon their oxygen isotope composition (Clayton, 2003). Although ureilites do not have a 1:1 match with any known chondrite parent body (Clayton and Mayeda, 1996), they plot along an extension of the CV field on the CCAM line, partly overlapping with oxygen isotope data for the CR chondrites (Weisberg et al., 2001). Recently, it has been shown that there are significant differences in the  $^{187}\text{Re}/^{188}\text{Os}$  and  $^{187}\text{Os}/^{188}\text{Os}$  of carbonaceous chondrites compared with ordinary and enstatite chondrites (Walker et al., 2002). There-

fore, the Os isotopic compositions may prove useful for ‘fingerprinting’ the provenance of planetesimals.

The Re-Os system is further explored in order to determine whether Re-Os chronology can provide additional constraints on the timing of ureilite differentiation. Available Sm-Nd and U-Pb ages for monomict ureilites are compatible with an early ( $\sim 4.56$  Ga) differentiation of the ureilite parent body (Goodrich and Lugmair, 1995; Torigoye-Kita et al., 1995b; Torigoye-Kita et al., 1995c). A Pb-Pb age ( $4.56 \pm 0.03$  Ga) consistent with this was obtained from apatite from the polymict ureilite DaG 319 (Kita et al., 2002). Preliminary  $^{182}\text{Hf}$ - $^{182}\text{W}$  data for 8 monomict ureilites suggests an old age for these ureilites comparable to the timescales for differentiation of the howardite-eucrite-diogenite parent body, i.e. within a few million years since solar system formation (Lee et al., 2005). More recent  $^{26}\text{Al}$ - $^{26}\text{Mg}$  data for the ureilites suggests extremely ancient ages from potentially older than the calcium-aluminum-rich inclusions (CAIs – commonly thought to represent the oldest solids that formed in our solar system) to about 0.5 Ma after CAI formation (Baker and Bizzarro, 2005). Polymict ureilites also contain Mn- and Al-rich clasts that yield ages of  $-4.5 \pm 0.4$  Ma relative to angrites using the  $^{53}\text{Mn}$ - $^{53}\text{Cr}$  short-lived radionuclide system (Goodrich et al., 2002), and  $\sim 5$  Ma after formation of CAIs using the  $^{26}\text{Al}$ - $^{26}\text{Mg}$  system (Kita et al., 2003). Both methods yield a similar absolute age of  $\sim 4.562$  Ga for these clasts, and hence, assuming that these clasts represent indigenous ureilitic lithologies, an estimate for the last magmatic activity on the ureilite parent body. However, some subsequent isotopic disturbance of ureilites is indicated by the most recent times of Ar degassing that have been inferred as 4.5 to 4.6 Ga for PCA 82506, but 4.1 Ga for Kenna and 3.3 to 3.7 Ga for Novo-Urei (Bogard and Garrison,

1994). In addition, it has been suggested for Kenna and Novo-Urei that material enriched in the light rare earth elements (LREE) was mobilized on the ureilite parent body at that time, possibly by impact (Goodrich and Lugmair, 1995; Goodrich et al., 1995), although it has also been argued that the 3.79 Ga Sm-Nd isochron for the ureilites Kenna, Novo-Urei, ALHA 77257 and Goalpara represents a mixing line due to terrestrial contamination (Torigoye-Kita et al., 1995a, 1995b).

Natural Re is comprised of two isotopes,  $^{185}\text{Re}$  (37.40 atom%) and  $^{187}\text{Re}$  (62.60 atom%). The isotope  $^{187}\text{Re}$  decays by  $\beta^-$  emission to  $^{187}\text{Os}$  with a half-life of 41.6 Ga (Shen et al., 1996; Smoliar et al., 1996). Within the past 20 years, the  $^{187}\text{Re}$ - $^{187}\text{Os}$  system has established its rank among the radiogenic isotope systems commonly applied to cosmochemical issues (Shirey and Walker, 1998). Because Re and Os are both refractory and siderophile elements, the Re-Os system has proven useful in studies of primitive solar system materials such as the iron meteorites (Shen et al., 1996; Smoliar et al., 1996), but also CAIs and chondrites (Becker et al., 2001; Brandon et al., 2005a; Brandon et al., 2005b; Walker et al., 2002). Because the ureilites contain significant amounts of metal, a Re-Os study seems appropriate to shed some light on their formation history.

## 2. Samples

Twenty-two ureilites were analyzed in this study, including 16 Antarctic samples provided by the Astromaterials and Research Exploration Science Directorate at NASA-JSC (Table 1). Dyalpur (Field Museum of Natural History, Chicago), Jalanash (Institut für Planetologie, Münster) and Novo-Urei (National Museum of Natural History, Washington, USNM 2969) are observed falls from India, Mongolia and Russia, respectively.

Goalpara (USNM 5992) and Kenna (USNM 5825) were found in India and the US, respectively. DaG 319, found in the Libyan Desert, was purchased for this study. The sample suite spans a broad compositional range representative of the ureilite group as a whole. Most of the ureilites in this study have typical monomict olivine-pigeonite lithologies. In the initial thin section description of EET 96042 no pyroxene was observed (Satterwhite and Lindstrom, 1998), whereas another study lists this ureilite as pigeonite-bearing (Cloutis and Hudon, 2004). MET 01085 has been described as olivine-free (Satterwhite and Allen, 2002); it is however, like all ureilites, sufficiently coarse-grained to provide the possibility that other sections of MET 01085 do contain olivine. LEW 85440, META 78008 and ALH 82130 (paired with ALH 82106, not analyzed) are augite-bearing. LEW 88774 is augite-bearing and also notable for being one of only two ureilites known to contain primary chromite. DaG 319 is the only polymict ureilite analyzed in this study. Weathering categories A, B, and C (Table 1) are used by the Meteorite Working Group at the NASA Johnson Space Center in Houston for Antarctic meteorite finds, denoting minor, moderate, and severe rustiness of hand specimens. Weathering levels of the analyzed ureilites vary from very fresh (including the falls) to severely rusty (LEW 86216). The warm-desert sample DaG 319 is classified as W2, meaning moderate oxidation of metal, 20-60% being affected (Wlotzka, 1993). Shock levels for the ureilites are taken from the compilation given in Mittlefehldt et al. (1998).

### **3. Analytical techniques**

All sample preparation and chemistry was performed at the Johnson Space Center. Initial sample masses ranged from 75 mg (LEW 86216) to 5.2 g (GRA 98032). Most



of the ureilite samples were coarsely crushed to mm-sized pieces using a ceramic alumina mortar and pestle. Three of the largest samples (DaG 319: 5g; Goalpara: 3g; Kenna: 4.5g) were crushed to a fine powder. Between samples, the mortar and pestle were cleaned by grinding quartz sand in ethanol to a fine powder, and then boiled in hot dilute nitric acid, followed by multiple rinses with Milli-Q ultra-pure water. Approximately 20 to 30 mg of rock chips (usually in 1 to 4 pieces) were handpicked from the coarsely crushed samples. If the outer rind of the meteorite was part of the original sample chip, material from the interior was taken for analyses. Leaching of the samples in dilute acids to reduce the effects of weathering in the Antarctic environment was not performed, because this could remove iron oxides (which likely contain Re and Os) in more strongly weathered samples. One sample (LEW 86216) showed very strong alteration in the form of pervasive iron oxides throughout the sample. In this case, the optically freshest parts were used for analysis, biasing the LEW 86216 analysis towards the iron oxide-poor side. To investigate sample reproducibility, sample GRA 98032 (total sample mass 5.2 g, weathering level C) was analyzed three times. A subsample of 90 mg rock chips was ground to a fine powder. Two ~20 mg subsamples of GRA 98032 were taken from this 90 mg powdered aliquot, another ~20 mg subsample was taken from the remaining rock chips (Table 1).

Sample digestion followed established procedures (Shirey and Walker, 1995). The rock chips or powders (~20 mg) were loaded in quartz Carius tubes, followed by addition of 2 ml concentrated HCl and a mixed spike enriched in  $^{185}\text{Re}$  and  $^{190}\text{Os}$  (Brandon et al., 2005a). The mixtures were then frozen in a dry-ice/ethanol slush before adding 4 ml of concentrated  $\text{HNO}_3$  resulting in a reverse aqua regia solution. The Carius tubes were then sealed and heated in an oven for 72 hours at 230°C. The Carius tube digestion

method is effective at dissolving metal and reaching spike-sample equilibration (Walker et al., 2002), but it is not a total dissolution method for silicate-bearing rocks. However, olivine and pyroxene are strongly attacked during the digestion procedure and the ureilite silicate rock matrix completely disintegrated. Moreover, the bulk of ureilite HSE are hosted in interstitial metal phases (Janssens et al., 1987) and the high-temperature igneous processing of ureilites precludes the occurrence of highly refractory primordial HSE-bearing alloys as observed for example in CAIs (Becker et al., 2001), or other refractory phases as observed in unequilibrated bulk chondrites (Brandon et al., 2005b). After digestion, Os was extracted from the reverse aqua regia using carbon tetrachloride solvent, then back extracted into HBr (Cohen and Waters, 1996), and finally purified by microdistillation (Birck et al., 1997). Following Os extraction, the aqua regia solutions were dried down and converted into chloride form, dissolved in 0.15N HCl, and loaded onto cation exchange columns for Re separation and purification (Puchtel and Humayun, 2001).

Osmium cuts were loaded onto Pt filaments with a mixed NaOH-Ba(OH)<sub>2</sub> emitter. The concentrations and isotopic compositions reported in Table 1 were measured at the Johnson Space Center by negative ion thermal ionization mass spectrometry (N-TIMS) using a Thermo Finnigan Triton in static multicollection mode. Oxygen corrections were made using the oxygen isotope composition measured on 2 ng loads of ReO<sub>4</sub><sup>-</sup> on the Faraday cups. The O<sub>2</sub> pressures in the source were maintained in the range of 1-3×10<sup>-7</sup> mbar for all runs. The <sup>187</sup>Re interference on <sup>187</sup>Os was monitored at mass 233 (<sup>185</sup>Re<sup>16</sup>O<sub>3</sub><sup>-</sup>) and was not observed during any of the sample measurements. After oxygen corrections were performed on the raw data, instrumental mass fractionation was corrected using <sup>192</sup>Os/<sup>188</sup>Os = 3.083 and the exponential law. The Johnson-Matthey Os iso-

tope standard gave  $^{187}\text{Os}/^{188}\text{Os} = 0.11380 \pm 1$  ( $2\sigma$ ) during the analytical period, identical to previously published values (Brandon et al., 2005a; Brandon et al., 2005b; Brandon et al., 1999).

The Re isotopic compositions were measured on a Finnigan Element™ single-collector, magnetic sector, high-resolution ICP-MS at the National High Magnetic Field Laboratory using an ESI™ PFA spray chamber and an ESI™ 100 $\mu\text{l}/\text{min}$  nebulizer (Puchtel and Humayun, 2001). The samples were run alternately with matrix solution blanks and a standard solution of similar Re concentration to the sample solutions. Instrumental mass fractionation was corrected by normalizing the measured  $^{185}\text{Re}/^{187}\text{Re}$  used for isotope dilution (ID) calculations with a mass fractionation factor obtained by comparing the measured  $^{185}\text{Re}/^{187}\text{Re}$  in the standard solution to the natural value (0.5974). For the majority of samples, the internal precision of individual ID-ICP-MS runs for the determination of Re concentrations (Table 1) had uncertainties of  $0.54 \pm 0.34\%$  ( $2\sigma$ ,  $n=25$ , Table 1). For 5 samples (Dyalpur, Goalpara, LEW 88774, LEW 85440, DaG 319) the Re chemistry yields were low, resulting in larger errors on  $^{187}\text{Re}/^{185}\text{Re}$  of 1.2 to 2.4% (Table 1). Total external uncertainties ( $2\sigma$ ) for Os concentrations from ID-TIMS analyses are estimated to be better than 0.1%. The total analytical blanks ranged between 2 and 22 (average 8) pg Re, and between 0.2 to 3.7 (average 1.4) pg Os. Blank corrections on sample Re concentrations averaged 2%, but were 11 and 18% for the samples LEW 85328 and ALHA 81101 that have very low Re abundances. Blank corrections on sample Os concentrations were  $<0.1\%$ . Because samples ALHA 81101 and LEW 85328 were also largely overspiked during a first set of measurements, these samples were subsequently duplicated along with 2 other samples from a fresh dis-

solution of a larger (40-70 mg) subsample of the coarse rock chips left from initial sample crushing (i.e. ALHA 81101, EET 96042, GRA 95205, and LEW 85328, cf. Table 1). The spiked Re cuts of three additional duplicates of homogenized sample powders (DaG 319, Goalpara, and Kenna) have been split for parallel N-TIMS determinations. Because N-TIMS analyses resulted in slightly lower uncertainties of calculated Re concentrations, these data are reported for the DaG 319, Goalpara and Kenna duplicates in Table 1.

#### **4. Results**

Measured Re abundances for the studied ureilites range from  $\sim 0.1$  times CI (Horan et al., 2003) in ALHA 81101 to  $\sim 2$  times CI in ALHA 78019, whereas Os abundances in the same ureilites range from  $\sim 0.1$  times CI to  $\sim 2.6$  times CI (Table 1). Compared to previous Re-Os data from neutron activation analysis for the same ureilites (Warren et al., 2006), the Re concentrations deviate between -60% (ALHA 81101) to +48% (PCA 82506), and Os deviates from -34% (Novo-Urei) to +164% (META 78008) from the literature values, respectively. Sample sizes in previous studies ranged between 100 to 500 mg, whereas sample sizes in this study ranged mostly between 20 and 30 mg (Table 1), but also included three large (3 to 5 grams) homogenized samples. It is not the intention of this study to determine absolute Re and Os abundances in ureilites, and 20 mg fractions of ureilites may not be representative of the whole meteorites. Even Kenna (4.5 grams of homogenized powder) differs from the literature Os concentration value (Warren et al., 2006) by 30% in this study. Considering that allocated sample masses are usually much smaller than 4.5 grams, and that many ureilite meteorites have less than 4.5

grams of total mass, it seems that absolute trace element abundances in ureilites will generally remain afflicted with considerable uncertainty.

The external reproducibilities of Re and Os concentration measurements for 20 mg aliquots taken from the large (3 to 5 grams) powdered samples Goalpara, DaG 319 and Kenna are within 1.1 and 1.9%, 4.6 and 3.5%, and 5.7 and 5.6%, respectively, and decrease with overall increasing Re-Os abundances in the samples (Fig. 1a). The reproducibility of the Re/Os ratio, however, increases with increasing Re-Os abundances, and is within 2.0, 0.6 and 0.2% for these samples, respectively. Three subsamples (powders and chips) of sample GRA 98032 showed virtually identical Os concentrations, but large Re abundance variations (Fig. 1a). Duplicate analyses of individual rock chip subsamples show more scatter with no obvious dependence on absolute Re-Os abundances, or weathering level (Fig. 1b).

The Os isotope analyses of all analyzed ureilites, including 9 duplicates, are reported in Table 1. The ureilite  $^{187}\text{Os}/^{188}\text{Os}$  form a unimodal distribution, and range from 0.11739 to 0.13018, with a mean of  $0.1258 \pm 0.0023$  ( $1\sigma$ ) (Fig. 2). The polymict ureilite DaG 319, which contains feldspathic lithologies that may be derived from the missing silicate melts from the ureilite parent body (Cohen et al., 2004; Downes and Mittlefehldt, 2006; Goodrich et al., 2004; Ikeda and Prinz, 2001; Kita et al., 2004), has  $^{187}\text{Os}/^{188}\text{Os}$  of  $0.1262 \pm 0.0004$  ( $2\sigma$ ) indistinguishable from the monomict ureilite average [ $0.1257 \pm 0.0024$  ( $1\sigma$ )]. Inverse Os concentrations and Os isotopic compositions of the ureilites are ( $r^2=0.66$ ) correlated (Fig. 3A). There are, however, no significant correlations of Os isotopic compositions with pyroxene (i.e. ureilite) type, olivine (Fig. 3B) or pyroxene (not

shown) core compositions, shock level (Fig. 3C), modal pyroxene content,  $\delta^{33}\text{S}$  (Fig. 3D), or  $\Delta^{17}\text{O}$  (Fig. 3E).

## **5. Discussion**

### **5.1. Effects of terrestrial weathering on ureilite Re and Os abundances and Os isotopes**

As pointed out in earlier work (Brandon et al., 2000; Esser and Turekian, 1993; Peucker-Ehrenbrink and Blum, 1998; Walker et al., 2002), terrestrial crustal materials typically have  $^{187}\text{Os}/^{188}\text{Os}$  of  $\geq 1$ , and widely varying, very high Re/Os. However, because of the relatively high concentrations of Os in the ureilite samples of 50 – 1200 ppb (Table 1), combined with the low concentrations of Os in crustal rocks and minerals of  $<0.05$  ppb, i.e. 1000 to 20000 times less Os compared to the samples, it is highly unlikely that  $^{187}\text{Os}/^{188}\text{Os}$  ratios were changed by Antarctic weathering. [This is also consistent with a recent study on the weathering behavior of CK carbonaceous chondrites \(Huber et al., 2006\) where even severely weathered Antarctic samples showed no resolvable anomaly in their Os abundance compared to other CK finds and falls.](#) Re was not measured in the latter study; it is, however, expected to be more mobile during terrestrial weathering than other platinum group elements (Jaffe et al., 2002). The main Re and Os-bearing phase in ureilites is kamacite that is associated with carbonaceous material (Janssens et al., 1987; Wänke et al., 1972; Wlotzka, 1972). Fractionations in whole rock Re/Os should occur only if Re and Os, once weathered out of the metal via formation of hydrous Fe-oxides, show largely different mobility within their host rock.

To investigate the effects of weathering on Re and Os in the studied suite of ureilites, time averaged  $^{187}\text{Re}/^{188}\text{Os}$  were calculated from the measured  $^{187}\text{Os}/^{188}\text{Os}$  of the samples (Table 1) assuming a formation age of the ureilites of 4562 Ma (Goodrich et al., 2002; Kita et al., 2003; Torigoye-Kita et al., 1995c) and closed-system Re/Os evolution. From these ratios, and the measured Os concentrations, average time-integrated Re concentrations were calculated for each ureilite (Table 1), and are compared with the measured Re concentrations (Fig. 4). The measured and calculated Re concentrations are correlated and scatter around a 1:1 reference line (Fig. 4). The visible evaporite minerals reported for two ureilites (weathering grade ‘e’, Table 1) do not seem to additionally disturb the Re-Os system. Correlation coefficients are similarly high among groups of samples classified as weathering grades A and B with  $r^2 = 0.91$  and  $0.94$ , respectively. In contrast, two samples classified as C (LEW 86216 and three duplicates of GRA 98032, Table 1) show large scatter ( $r^2 = 0.31$ ). Two independent subsamples of GRA 98032 (chips and powder coming from different parts of the original sample slab) have consistent Os abundances within 3.5% of each other, but variable Re abundances (Fig. 1A), which can be interpreted as Re mobility, but Os immobility in the hydrous Antarctic environment. Furthermore, the discrepancy in Re concentration between the two powdered subsamples of GRA 98032 suggests that homogenization was not achieved during sample grinding for this sample. Re mobility is also indicated for the most strongly weathered sample LEW 86216 that shows the largest Re loss (-72%) (Fig. 4). Because pieces of LEW 86216 that showed major rustiness were avoided for analyses, it is concluded that Re was either preferentially transferred to the iron oxides (short-range redistribution), or removed from the sample (long-range redistribution). Long-range redistribution of Re is more consistent

with the general Re loss indicated for the large homogenized sample Kenna (-19%) (Fig. 1A). The ureilites with intermediate and minor weathering grades span a range of measured Re concentrations from approximately -40% to +40% relative to their calculated Re concentrations (Fig. 4, Table 1). In comparison, the warm-desert sample DaG 319 (weathering level W2) lost 44% of its Re when compared to the calculated Re concentration from Os isotopes (Fig. 4).

The observed variability of Os concentrations in duplicate analyses of independent subsamples of ureilite rock chip (GRA 95205 and EET 96042, Fig. 1), and the much higher Os concentrations of samples GRA 95205 (+76%) and META 78008 (+164%) compared to previous data (Warren et al., 2006), are attributed to heterogeneous distribution of HSE-bearing metals within individual ureilite subsamples. A similar conclusion was made in studies of Os isotope/HSE systematics in chondrites (Horan et al., 2003; Walker et al., 2002). Very few Fe-metal abundances are reported for ureilites (Jarosewich, 1990; Wiik, 1972; Yanai and Noda, 2005). However, because of their variable distribution on the centimeter scale, Fe-metal abundances in a specific ureilite likely depend on the specific aliquot or thin section analyzed. In conclusion, absolute abundances of Os appear to be affected by heterogeneous distribution of metal, while those of Re have additionally been affected by weathering processes, both of which reduce the precision with which conclusions can be drawn from absolute Re and Os abundances about HSE distribution during ureilite petrogenesis. However, Os isotopes should be much less affected by such sample heterogeneity because even if metal was mobile in the ureilite parent body, the Re/Os and hence present-day  $^{187}\text{Os}/^{188}\text{Os}$  isotopic composition of that metal should be fairly homogeneous.



When plotted in a  $^{187}\text{Os}/^{188}\text{Os}$  versus  $^{187}\text{Re}/^{188}\text{Os}$  isochron diagram (Fig. 5A), most of the data scatter around a horizontal zero age isochron, consistent with recent mobilization and open system behavior of Re. There is no indication that  $^{187}\text{Re}/^{188}\text{Os}$  or  $^{187}\text{Os}/^{188}\text{Os}$  ratios of strongly weathered samples (weathering grade C) were preferentially shifted to superchondritic values, as would be expected from terrestrial contaminants. Because of the overall large scatter in ureilite Re/Os, however, no meaningful age information can be deduced from the whole rock data. Even when the dataset is restricted to the group of ureilite falls, no useful isochron can be fitted to the data because of the insufficient range in Re/Os. However, a subset of the ureilite whole rock data that includes the falls clusters more closely around the 4558 Ma group IIIA iron meteorite reference isochron (Fig. 5A), and two of the three ureilite falls studied have a measured Re-Os isotope composition that is indistinguishable from that of carbonaceous chondrites (Walker et al., 2002). Argon isotopic data for the Kenna and Novo-Urei ureilites suggest disturbances of at  $\sim 4.1$  Ga and 3.3 to 3.7 Ga (Bogard and Garrison, 1994), respectively, consistent with the suggested time of LREE-mobilization in these meteorites at  $\sim 3.79$  Ga (Goodrich and Lugmair, 1995; Goodrich et al., 1991). Although the Re-Os data for Kenna is compromised by too large weathering effects for a reliable Re-Os age estimate, the fresh ureilite fall Novo-Urei plots close to the 4558 Ma reference isochron (Fig. 5B). Thus, whatever caused the argon isotopic disturbances in Novo-Urei did not produce a noticeable effect on the Re-Os isotope system.

## **5.2. The provenance of the ureilite parent body**

### **5.2.1. Ureilite HSE systematics**

Ureilite HSE budgets have been suggested to be a mixture of a refractory-rich component that is associated with the veins, and a refractory-poor component associated with the silicates (Boynton et al., 1976; Higuchi et al., 1976; Janssens et al., 1987; Spitz and Boynton, 1991; Wänke et al., 1972). Later work has shown that ureilite HSE abundance patterns are consistent with solid metal/liquid metal fractionation, indicating that residual metal is present in ureilites (Goodrich et al., 1987a; Goodrich et al., 1987b; Humayun et al., 2005; Janssens et al., 1987; Jones and Goodrich, 1989; Warren and Huber, 2006; Warren et al., 2006). A good match for the siderophile element patterns of ureilites with high Re and Os abundances is obtained assuming that ureilites are residues of carbonaceous chondrite precursors after extraction of a S-rich (>15%) liquid metal component (Humayun et al., 2005). Osmium isotopes provide a means to test the hypothesis that these two components with low versus high Re and Os abundances are characterized by distinct Os isotopic compositions. Mixing of two isotopically distinct Os components should form a straight line in a  $1/\text{Os}$  versus  $^{187}\text{Os}/^{188}\text{Os}$  plot (Fig. 3B). Although the observed correlation is defined mainly by the two depleted samples ALHA 81101 and MET 01085 (Fig. 3A), elimination of these two samples in the calculation still yields a significant correlation (95.1% confidence) in the remaining data set. It can therefore be concluded that Os isotopes are consistent with mixing of an Os-poor component (probably associated with the silicates) with long-term depletion in Re/Os, and Os-rich residual metal that is associated with the carbonaceous material and has more carbonaceous chondrite-like Re/Os and consequent  $^{187}\text{Os}/^{188}\text{Os}$  of  $\sim 0.1265$  (Fig. 3A).

The mean and distribution of  $^{187}\text{Os}/^{188}\text{Os}$  measured for the ureilites ( $0.1258 \pm 0.0023$ ) are similar to those measured in carbonaceous chondrites ( $0.1258 \pm 0.0018$ ) and

in CAIs with relatively unfractionated REE patterns ( $0.1265 \pm 0.0003$ ), but different from those measured for ordinary chondrites ( $0.1281 \pm 0.0020$ ) or enstatite chondrites ( $0.1281 \pm 0.0005$ ). The polymict (brecciated) ureilites are composed of >98% material similar to the monomict ureilites (Downes and Mittlefehldt, 2006; Goodrich et al., 2004). These materials encompass an identical range of mineralogical compositions to that shown by all known monomict ureilites, and also show similar relative proportions of the different petrologic types. The  $^{187}\text{Os}/^{188}\text{Os}$  of  $0.1262 \pm 0.0004$  ( $2\sigma$ , ext.) measured in this study for the polymict ureilite DaG 319 is consistent with the average  $^{187}\text{Os}/^{188}\text{Os}$  of the 21 monomict ureilites, and thus in agreement with the petrographic observations.

The division of bulk chondrites on the basis of  $^{187}\text{Os}/^{188}\text{Os}$  into two groups (carbonaceous versus ordinary/enstatite chondrites) may result from differential high-temperature nebular condensation of Re and Os into refractory element-bearing alloys and their subsequent isolation and incorporation into the precursor materials of certain chondrite groups (Walker et al., 2002). Alternatively, the observed disparate Re/Os could also be a result of lower-temperature oxidation or heating loss of Re in the precursor materials of carbonaceous chondrite components (Walker et al., 2002). The overlapping  $^{187}\text{Os}/^{188}\text{Os}$  for certain CAIs and carbonaceous chondrites, however, suggest a bulk solar system value of  $\sim 0.1265$  (Becker et al., 2001), and the high-temperature condensation scenario therefore seems more plausible. In the following, two conclusions are derived from the similar distribution and mean of  $^{187}\text{Os}/^{188}\text{Os}$  measured for ureilites and for carbonaceous chondrites (Fig. 2).

First, the similarly narrow range of  $^{187}\text{Os}/^{188}\text{Os}$  (i.e. similar standard deviation) observed for ureilites and carbonaceous chondrites is taken as evidence that partial melt-

ing in the ureilite parent body, accumulation of metal, and different degrees of shock (Fig. 3C) early in ureilite petrogenesis did not lead to a significant fractionation of Re from Os. In particular, metallic iron (i.e. residual metal) must have been a stable phase in the early fractionation history, because in the absence of a metal phase, Re should behave incompatibly and may have been lost along with the missing ureilitic melt component. For example, if the ureilite precursor material experienced early magmatic processing above the iron-wuestite oxygen buffer, Re and Os partitioning would be expected to be comparable to melt extraction in the terrestrial upper mantle at similar oxygen fugacities. For example,  $\text{Al}_2\text{O}_3$  contents correlate with  $^{187}\text{Os}/^{188}\text{Os}$  in terrestrial orogenic lherzolites (Reisberg and Lorand, 1995). This relationship is consistent with similar incompatibilities and proportional removal of  $\text{Al}_2\text{O}_3$  and Re from the peridotite residues during partial melting in an oxidized, metal-free terrestrial upper mantle. Terrestrial peridotite xenoliths from the non-convecting subcontinental lithospheric mantle have lost on average 63% of their Re compared to the present fertile convecting mantle (Shirey and Walker, 1998). Compared to average CI and CV chondrites (Wasson and Kallemeyn, 1988), average ureilites (Warren et al., 2006) have lost 66 and 83% of their aluminum, respectively, similar to the terrestrial analogue. Assuming a chondritic starting composition at 4562 Ma and a relative proportion of Re equivalent to  $\text{Al}_2\text{O}_3$  was lost (i.e. a present-day  $^{187}\text{Re}/^{188}\text{Os}$  ranging between 0.0666 and 0.1333), then the present-day average  $^{187}\text{Os}/^{188}\text{Os}$  for the ureilites should be  $\leq 0.1058$ . Because such low  $^{187}\text{Os}/^{188}\text{Os}$  is not observed for the ureilites studied here, it is concluded that Re was not significantly lost relative to  $\text{Al}_2\text{O}_3$  during melt extraction, and thus metal must have been a stable phase throughout ureilite magmatic processing. The presence of metal implies oxygen fugaci-

ties that were below the iron-wuestite oxygen buffer throughout the melting interval. It must be noted that some carbonaceous chondrites contain very little metal, and HSE may be concentrated in sulfides. Unlike ureilites, these carbonaceous chondrites did not undergo magmatic processing and melt removal, and therefore retained their bulk Re/Os.

Second, the similar mean of  $^{187}\text{Os}/^{188}\text{Os}$  measured for the ureilites and carbonaceous chondrites suggests that the ureilite parent body probably formed within the same region of the solar nebula as carbonaceous chondrites. This conclusion from Os isotopes is supported by oxygen isotopes of ureilites (Clayton and Mayeda, 1996). In the oxygen three-isotope plot (Fig. 6), ureilites plot along an extension of the CCAM, but are clearly distinct from ordinary and enstatite chondrites. The reason for the slightly lower-than-average  $^{187}\text{Os}/^{188}\text{Os}$  of ALHA 81101 and MET 01085 is unclear. While ALHA 81101 is a relatively unweathered, highly shocked, low forsterite (Fo), and pigeonite-poor ureilite characterized by the lowest overall Re and Os abundances measured in this study; MET 01085 is an unusual orthopyroxene-bearing ureilite with intermediate values for weathering grade, Re and Os abundances, Fo contents, and shock level (Table 1, Fig. 3A-C). Thus the low  $^{187}\text{Os}/^{188}\text{Os}$  can not be connected to any other petrologic feature, arguing against the presence of a distinct subchondritic  $^{187}\text{Os}/^{188}\text{Os}$  and Os-poor component associated with the silicates as suggested from the correlation in Figure 3A.

Subchondritic  $^{187}\text{Os}/^{188}\text{Os}$  could in principle result from extraction of a S-rich metal phase from the ureilites. Solid metal/liquid metal distribution coefficients ( $D^{\text{SM}/\text{LM}}$ ) for Re and Os during such liquid metal extraction are dependent on the S-content of the liquid metal phase (Chabot et al., 2003; Chabot and Jones, 2003; Rushmer et al., 2005), which decreases during batch melting as a function of temperature from the eutectic

composition (~30 wt% S in the liquid phase). Using the  $D^{\text{SM}}_{\text{LM}}$  parameterization of (Chabot and Jones, 2003), and assuming a simplified binary Fe-S system,  $D^{\text{SM}}_{\text{LM}}$  for Re and Os are in the order of several thousands at the Fe-S eutectic, but decrease to 34.9 and 36.9, respectively, for a liquid metal S-content of ~20 wt% that correspond to the peak metamorphic temperatures for the ureilites of 1300°C. The liquid fraction F produced at the Fe-S eutectic is dependent on the initial FeS to Fe-metal ratio of the ureilite protolith. However, assuming that the melt fraction was smaller than 90%, the calculated Re/Os fractionations in residual metal are smaller than 1.14%; and a fractionation in Re/Os at 4562 Ma of this magnitude would lead to a present-day deficit in  $^{187}\text{Os}/^{188}\text{Os}$  of the residual metal of 0.28%. It is concluded that extraction of a Fe-S liquid from the ureilites is unsuitable to explain the  $^{187}\text{Os}/^{188}\text{Os}$  deficits of ALHA 81101 and MET 01085 of 6.6 and 3.7%, respectively, relative to the ureilite average (Table 1).

A low  $^{187}\text{Os}/^{188}\text{Os}$  value [ $0.11980 \pm 0.00004$  ( $2\sigma$ )] similar to the two anomalous samples ALHA 81101 and MET 01085 has been measured for the undifferentiated CK4 chondrite Karoonda (Walker et al., 2002). This opens the possibility that the low  $^{187}\text{Os}/^{188}\text{Os}$  of ALHA 81101 and MET 01085 might reflect heterogeneous carbonaceous chondritic precursor material.

### **5.2.2. What is the ureilite precursor material?**

Ureilite Os isotope data are indistinguishable from those of carbonaceous chondrites, but a more specific assignment to a specific chondrite group is not possible with Os isotopes alone. In oxygen isotopes, ureilites do not have a 1:1 match with any carbonaceous chondrite parent body (Fig. 6). The ureilites plot along an extension of the CV

field on the CCAM line, partly overlapping with oxygen data for the CR chondrites (Weisberg et al., 2001). The CCAM line may be explained as a mixing line between  $^{16}\text{O}$ -rich condensates from the unaltered primary nebular gas having approximately -45‰ for both  $\delta^{17}\text{O}$  and  $\delta^{18}\text{O}$ , and material from the inner part of the accretion disk enriched in  $^{17}\text{O}$  and  $^{18}\text{O}$  by the photolysis of CO (Clayton, 2002). The large range in  $\delta^{18}\text{O}$  of each carbonaceous chondrite group compared to the ordinary and enstatite groups (Fig. 6) is probably a result of low-temperature aqueous alteration, which produces phyllosilicates enriched in the heavy isotopes (Clayton and Mayeda, 1999). For example, the phyllosilicate matrix of CM and CO chondrites is systematically enriched in  $\delta^{18}\text{O}$  and  $\delta^{17}\text{O}$  relative to the whole rock, and the tie-lines between whole-rock compositions and matrix compositions have slopes of  $\sim 0.7$ . In the case of the CV chondrites, the large range in  $\delta^{18}\text{O}$  may represent internal isotopic heterogeneity due to the presence of  $^{16}\text{O}$ -rich refractory phases (Clayton and Mayeda, 1999). The offset between CV chondrites and ureilites on the CCAM thus could be explained if the ureilite protolith is similar to CV chondrites, but contained less refractory phases. The close fit of the ureilites to the CCAM line with slope  $\sim 1$  can be taken to imply that aqueously altered carbonaceous chondrite material (CI, CM or CR) was at most a minor component of the ureilite parent body.

A CV-type precursor of the ureilites, however, is at odds with the carbon abundance in ureilites [0.7-6.6 wt%, average 2.9 wt% (Hudon et al., 2004; Mittlefehldt et al., 2005; Warren et al., 2006)], which is higher than that in CV chondrites by a factor of about 5, but comparable to CI and CR chondrites [averages 0.53, 3.45, and 2.0 wt%, respectively (Lodders and Fegley, 1998)]. The  $\delta^{13}\text{C}$  isotopic compositions of ureilites range from -11 to +1 ‰ (Hudon et al., 2004), and are also more akin to CI than CV chondrites

[ $\delta^{13}\text{C}$  ranging from -12 to -5 ‰ and -25 to -8 ‰, respectively (Grady and Wright, 2003)]. Although the carbon isotopic composition of the ureilites might have been altered by magmatic processing (Hudon et al., 2004), magmatic models are difficult to reconcile with the observed carbon enrichment of ureilites compared to CV chondrites. For example, loss of 30% silicate melt from the precursor would passively increase the carbon content of CV-type material to 0.8 wt%, far below the observed average abundance in ureilites. Chromium isotope data for Kenna and LEW 85440 suggest that the initial Cr isotope ratios of ureilites are distinct from those of carbonaceous chondrites, but similar to the HED parent body (Shukolyukov and Lugmair, 2006; Trinquier et al., 2005). However, the origin of  $^{54}\text{Cr}$  variation in the early solar system is as yet unclear and needs to be substantiated by further measurements. In conclusion, no known chondrite class seems to meet all the necessary conditions to provide a suitable ureilite precursor material. Nevertheless, oxygen and Os isotopes strongly suggest an affinity of ureilites to carbonaceous chondrites.

## **6. Conclusion**

In conclusion, absolute abundances of Os in ureilites appear to be affected by heterogeneous distribution of metal, while those of Re have additionally been affected by weathering processes. Because of Re losses during weathering, no meaningful age information can be deduced from the ureilite whole rock data. However, a subset of the ureilite whole rock data that includes the falls clusters more closely around the 4558 Ma group IIIA iron meteorite reference isochron, consistent with the old age of the ureilites obtained from other dating methods. From the similar mean and distribution of



$^{187}\text{Os}/^{188}\text{Os}$  measured for the ureilites and carbonaceous chondrites it is concluded that the ureilite parent body probably formed within the same region of the solar nebula as carbonaceous chondrites. Partial melting in the ureilite parent body, accumulation of metal, and different degrees of shock early in ureilite petrogenesis did not lead to a significant fractionation of Re from Os. In particular, metallic iron (i.e. residual metal) must have been a stable phase throughout the early ureilite magmatic history.

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