In search of late-stage planetary building blocks

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Abstract

Genetic contributions to the final stages of planetary growth, including materials associated with the giant Moon-formation impact, late accretion, and late heavy bombardment are examined using siderophile elements. Isotopic similarities between the Earth and Moon for both lithophile and siderophile elements collectively lead to the suggestion that the genetics of the building blocks for Earth, and the impactor involved in the Moon-forming event were broadly similar, and shared some strong genetic affinities with enstatite chondrites. The bulk genetic fingerprint of materials subsequently added to Earth by late accretion, defined as the addition of ~0.5 wt.% of Earth’s mass to the mantle, following cessation of core formation, was characterized by 187Os/188Os and Pd/Ir ratios that were also similar to those in some enstatite chondrites. However, the integrated fingerprint of late accreted matter differs from enstatite chondrites in terms of the relative abundances of certain other HSE, most notably Ru/Ir. The final ≤ 0.05 wt.% addition of material to the Earth and Moon, believed by some to be part of a late heavy bombardment, included a component with much more fractionated relative HSE abundances than evidenced in the average late accretionary component. Heterogeneous 182W/184W isotopic compositions of some ancient terrestrial rocks suggest that some very early-formed mantle domains remained chemically distinct for long periods of time following primary planetary accretion. This evidence for sluggish mixing of the early mantle suggests that if late accretionary contributions to the mantle were genetically diverse, it may be possible to isotopically identify the disparate primordial components in the terrestrial rock record using the siderophile element tracers Ru and Mo.

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1. Introduction

The origins of the rocky planets, especially with regard to assembly processes and the chemical nature of their building blocks, have been the topic of intense interest and debate for decades. It is now generally agreed that the terrestrial planets were built as a result of a three stage process, initially consisting of the growth of planetesimals from dust,
followed by the collisions of bodies of increasingly greater mass to create planetary embryos, culminating in the collision and accretion of the embryos to form planets during a giant impact stage of growth (e.g., Kokuba and Uda, 1998; Raymond et al., 2006; Morbidelli et al., 2012; Raymond et al., 2013). The final stage of major growth for the Earth likely occurred with the giant impact of a body comprising 5 wt.% or more of the mass of the present Earth, leading to the formation of the Moon (Hartmann and Davis, 1975; Canup and Asphaug, 2001; Canup, 2012; Ćuk and Stewart, 2012; Reuf et al., 2012). Minor accretion amounting to ~0.5 wt.% of the present Earth's mass continued after this event (Kimura et al., 1974; Chou, 1978).

In addition to constraining the dynamical processes involved in the construction of the rocky planets, it is equally important to assess the origins and chemical nature of the materials from which the planets were built. Comparisons of the bulk compositions of the terrestrial planets have commonly been made on the basis of chemical models developed for these planetary bodies, most notably the Earth, Moon, and Mars; that is, bodies from which we are reasonably confident we have samples. Such models often require the assumption that the bodies were constructed from a combination of materials that were compositionally similar to the primitive meteorites present in our collections (Wänke and Dreibus, 1988; Wänke et al., 1994; McDonough and Sun, 1995; Wänke, 2001; Taylor et al., 2006). The bulk planetary concentrations of a number of poorly-constrained elements have been estimated by applying bootstrapping methods which assume that the ratios of these elements to relatively well-constrained, geochemically-comparable major elements, are similar to the ratios observed in primitive meteorites (e.g., McDonough and Sun, 1995). Models of the chemical composition of inaccessible planetary reservoirs, such as Earth's core, often require these types of general assumptions (McDonough, 2003). Of course, it is unlikely that our meteorite collections sample the full compositional range of primitive materials involved with planetary accretion. Thus, some limitations in bootstrapping methods must be recognized. To circumvent this problem when assessing the compositions of some elements in inaccessible planetary reservoirs, such as cores, abundances can also be estimated by reverse modeling, using known or assumed mantle concentrations, and applying appropriate metal–silicate distribution coefficients. Models of this type, however, must make assumptions about the conditions of progressive metal–silicate segregation in a growing body (e.g., Rubie et al., 2011).

The chemical and genetic makeup of the rocky planets can potentially be further constrained by isotopic comparisons to one another and to primitive meteorites. For example, planetary materials exhibit a large range in mass independent variations in $\Delta^{17}$O ($\Delta$ per mili deviation in $^{17}$O/$^{16}$O from the terrestrial fractionation line), which can be used as genetic fingerprints of precursor materials. It has been hypothesized that the heterogeneities in $\Delta^{17}$O originated as a result of self-shielding effects in the photo-dissociation of CO by exposure to ultraviolet light within the solar nebula (e.g., Thiemens and Heidenreich, 1983; Clayton, 2002; Lyons and Young, 2005). Variations in $\Delta^{19}$O among differentiated bodies have, therefore, commonly been interpreted to reflect the formation of precursor materials at greater or lesser distances from the Sun, possibly coupled with time of formation (e.g., Yurimoto and Kuramoto, 2004). As an example of the application of O isotopes to issues of genetics, the similarity in the $\Delta^{17}$O of the Earth and enstatite chondrites has commonly been interpreted to indicate that the major building blocks of the Earth formed in a region of the protoplanetary disk similar to where enstatite chondrites formed (Clayton et al., 1984; Javoy et al., 2010). Conversely, differences in the $\Delta^{17}$O compositions of the Earth and Mars have been cited as evidence that the building blocks of these two bodies differed substantially (Franchi et al., 1999). Although there appears to be no perfect fit of all physical parameters between any types (or likely any combination of types) of primitive meteorites and the Earth, Moon or Mars, constraining the general categories of accretionary materials, nevertheless, remains an important objective of cosmochemistry.

Here, we focus mainly on the final ~10 to ~0.05 wt.% of Earth's accretion. Late stages of major terrestrial planetary accretion may have included the participation of materials that formed in different portions of the protoplanetary disk, and included water- and organic-rich materials (Weidenschilling et al., 1997; Chambers, 2001, 2004). Thus, although limited in mass, late stage planetary growth may have had a disproportionate effect on the volatile contents of the rocky planets (e.g., Kerridge, 1985; Balsiger et al., 1995; Eberhardt et al., 1995; Albarede et al., 2013; O'Brien et al., 2014; Rubie et al., 2015a). Further, late stage additions may have carried genetically distinct elemental and isotopic fingerprints. Here we use the term genetic to mean chemical and especially isotopic compositions of the primary nebular materials from which an object was built. Because of the comparatively limited mass contributed by these processes, elemental and isotopic tracers comprising major elements, such as O, are of limited value in constraining the nature of these final building blocks. Thus, we will instead explore the possibility of tracing the late-stages of planetary growth using insights gained from elemental and isotopic variability of so-called siderophile, or Fe-loving, elements.

In this overview, the elemental and isotopic fingerprints of late stage building blocks that may be recorded in mantle rocks from the Earth, as well as mantle-derived and impact generated rocks from Mars and the Moon, respectively, will be examined. In addition to considering the average elemental and isotopic characteristics of siderophile elements contained in the silicate portions of these bodies, we will also explore the possibility that the signals of individual building blocks might be identified through small differences in the isotopic compositions of the siderophile elements Ru and Mo, which varied among early solar system materials as a result of their incorporating differing proportions of diverse nucleosynthetic components. The basis for this optimism is the discovery that primordial mantle heterogeneities, recorded by lithophile, atmospheric and siderophile short-lived radiogenic isotope systems, survived long enough to be preserved in the terrestrial rock record (Caro et al., 2003; Willbold et al., 2011; Mukhopadhyay, 2012; Touboul et al., 2012, 2014). If the interpretations of long-lived chemical/isotopic heterogeneity in the mantle presented by these studies are correct, isotopically distinct domains within the mantle, imparted during late stage accretion of genetically distinct materials, might also be preserved in the rock record.

2. Overview of siderophile elements

Siderophile elements are those elements that strongly partition into liquid metallic Fe relative to silicate melt, and are consequently concentrated, to greater or lesser extents, in the cores of the rocky planets (Goldschmidt, 1937). Because of this, their concentrations in silicate mantles and crusts are low compared to primitive meteorites, the compositions of which are presumed to be representative of the majority of the planetesimals involved in the final stages of rocky planet accretion (Anders and Grevesse, 1988). Siderophile trace elements are commonly divided into sub-groups based on the intensity of their siderophile tendencies under the typical 1 atmosphere experimental conditions initially employed to characterize the nature of metal–silicate partitioning of these elements (e.g., Kimura et al., 1974; Borisov et al., 1994). The moderately siderophile elements (MSE), including W, Co, Ag, Ni, Ge, and Mo, are characterized by metal–silicate D values (concentration ratio of an element in liquid metal to liquid silicate) ranging from about 10 to 1000. The highly siderophile elements (HSE), including Re, Os, Ir, Ru, Pt, Rh, Au and Pd, are characterized by D values greater than 10,000. One important characteristic of siderophile elements is that the intensity of their siderophile behavior can shift considerably at increasingly higher temperatures and pressures (e.g., Ringwood, 1968; Murthy, 1991; Li and Agee, 1996, 2001; Holzheid et al., 2006). The general tendency of most, but not all, siderophile elements is towards lower D values, as pressure and temperature conditions increase (Rigter and Drake, 1997; Mann et al., 2012). Their partitioning characteristics are...
also affected by other intensive parameters of a given planetary body, such as oxygen fugacity (\(f_O^2\)), as well as the compositions of the participating metal and silicate phases (Wade and Wood, 2005; Cottrell and Walker, 2006; Wade et al., 2012). These shifts in partitioning behavior are important to recognize when considering issues of planetary growth and core formation. Depending upon the conditions at which metal last equilibrated with silicates, during progressive core formation in a growing body, appropriate D values likely changed considerably during the growth of sizable planetary bodies, thus, affecting the final absolute and relative concentrations of the siderophile elements in these mantles (e.g., Wade and Wood, 2005; Rubie et al., 2011).

For the purposes of genetic tracing of late stage building blocks, it is also important to recognize that the processes leading to the present abundances of the MSE contained within the silicate portions of the rocky planets may not have been the same as for the HSE. The chondrite-normalized abundances of the MSE estimated for the bulk silicate Earth (BSE) vary considerably (Fig. 1). Experimental studies have shown that the abundances of many of the MSE can be accounted for if metal–silicate equilibration occurred at elevated temperatures and pressures (e.g., Hillgren et al., 1994; Rühter et al., 1997; Li and Agee, 2001; Mann et al., 2009; Siebert et al., 2011; Wade et al., 2012). Thus, numerous studies have concluded that the abundances of these elements are consistent with metal–silicate equilibration at an average depth equivalent to pressures of 20–60 GPa. A major shift in \(f_O^2\) of the terrestrial mantle, resulting from either the disproportionation of ferrous iron into ferric iron plus metal, as occurs in Bridgmanite at high pressure (Frost et al., 2004; Frost and McCammon, 2008), or the partitioning of Si into core-forming metal, has also been proposed as having had a significant influence on the final concentrations of the MSE in the Earth's mantle (e.g., Wade and Wood, 2005; Rubie et al., 2011, 2015b).

In contrast to the MSE, the HSE occur in approximately chondritic relative proportions in the bulk silicate Earth (BSE) (Fig. 1) (Chou, 1978; Morgan, 1986; Meisel et al. 2001; Becker et al., 2006; Fischer-Gödde et al., 2011), and absolute abundances are only ~200 times lower than bulk CI chondrite abundances (Morgan, 1986). Further, the \(^{187}\text{Os/188}\text{Os}\) and \(^{186}\text{Os/188}\text{Os}\) ratios estimated for the BSE, that reflect the long-term decay of \(^{187}\text{Re}\) and \(^{190}\text{Pt}\) (where \(t_1/2\) for \(^{187}\text{Re}\) and \(^{190}\text{Pt}\) are ~42 and 450 Gyr, respectively), are also within the range of chondritic meteorites, thus providing a robust record for time-integrated chondritic Re/Os and Pt/Os (Morgan, 1985; Walker et al., 1997; Meisel et al., 2001; Brandon et al., 2006). These characteristics of the HSE in the BSE do not appear to be the consequence of high pressure and temperature metal–silicate equilibration. Experimental studies have shown that large differences in the metal–silicate distribution coefficients of siderophile elements at relevant temperatures and pressures would have led to non-chondritic absolute abundances in the mantle (Holzheid et al., 2000; Brenan and McDonough, 2009; Mann et al., 2012). This is observed for MSE but not for HSE (Fig. 1). Instead, the HSE may owe their presence in the BSE to continued accretion of planetesimals with bulk chondritic compositions, following core formation (Kimura et al., 1974; Chou, 1978). This process has been commonly termed the late meteorite influx, the late veneer, or late accretion. This terminology can be confusing because different authors have used the terms to mean different things. For example, some authors have used the terms “late veneer” and “late accretion” to describe all material accreted after the end of core formation, and all material accreted after the last giant impact, respectively (e.g., Jacobson et al., 2014; Morbidelli and Word, 2015). If the giant impact was the last event that led to growth of the core, then the terms are synonymous. The term late accretion has even been used to include the final major stage of terrestrial growth by giant impact (e.g., Schönbachler et al., 2010). Henceforth, here we will use the term late accretion to mean all additions following cessation of core formation.

Late accretion is a viable mechanism to explain HSE abundances in the BSE because the addition of bodies with chondritic bulk compositions to silicate mantles would have led to the establishment of comparatively high absolute, and chondritic relative abundances of HSE in the affected mantles. In the case of Earth, mass balance constraints suggest that it would be necessary to add ~0.5 wt.% of the mass of the total Earth (~2 \times 10^{22} \text{kg}) to the mantle by late accretion in order to account for present-day mantle HSE abundances (Walker, 2009). Implicit in such models are the assumptions that abundances of all HSE in the mantle were very low prior to late accretion, and that any metal present in late stage impactors was ultimately oxidized and incorporated in the mantle, so that the siderophile elements added to the mantle after core formation remained in the mantle (Kimura et al., 1974). The late accretion hypothesis is currently the only viable process to explain the HSE characteristics of the BSE (Walker, 2009). Nevertheless, the hypothesis is not without weaknesses. For example, the considerable apparent difference in HSE abundances in the BSE compared to the bulk silicate Moon is a potential major problem that will be discussed in Section 5.2.

Appealing to two distinct mechanisms for the establishment of siderophile elements in the terrestrial mantle is not necessarily a conflict. Models such as those proposed by Wade and Wood (2005) and Rubie et al. (2011, 2015a) would lead to the establishment of typically >90% of the MSE abundances in the mantle by high pressure and temperature metal–silicate partitioning. Repeated processing of metal through the mantle, as a result of growth by giant impacts and the resulting multiple stages of magma ocean formation and evolution, would lead to a biasing of the MSE present in the mantle today towards the mass added by the later major stages of accretion (Azbel et al., 1993; Kramers, 1998; Rubie et al., 2011, 2015b). For example, calculations using the progressive growth model of Kramers (1998) suggest that >80% of the W present in the mantle today was added by accretion of the final 10–20 wt.% of the Earth. By comparison, mass balance constraints suggest that late accretion of ~0.5 wt.% mass addition would add >95% of the HSE, but a maximum of only about 10% of an MSE, such as W, to the mantle total. Thus, as noted by Dauphas et al. (2004), the genetics of the MSE present in the mantle are not required to have been the same as the genetics of the HSE.

3. Isotopic tracers

Two types of isotopic tracers of building blocks are considered here; radiogenic and nucleosynthetic. Long-lived radiogenic tracers have long proved to be useful for constraining the vigor and nature of mixing in the silicate portions of planets. For example, long-lived systems including \(^{87}\text{Rb–87}\text{Sr}\), \(^{136}\text{Sm–142}\text{Nd}\), \(^{176}\text{Lu–176}\text{Hf}\), and \(^{238}\text{U–235}\text{U–232}\text{Th–206,207,208}\text{Pb}\) have been extensively used to trace mantle mixing over Earth history, particularly with respect to recycled crustal components (Zindler and Hart, 1986; Hofmann, 2003). Of the long-lived systems consisting of HSE, however, only the \(^{187}\text{Re–187}\text{Os}\) isotopic system, and to a lesser extent the \(^{186}\text{Pt–186}\text{Os}\) system, have been shown to be useful for characterizing late stages of planetary accretion to Earth, the Moon, and Mars.
Short-lived radiogenic isotopic systems have much greater utility for examining the timing of early planetary accretion and differentiation processes with high resolution (e.g., Caro et al., 2003; Boyet and Carlson, 2005; Foley et al., 2005; Debaille et al., 2013; McLeod et al., 2014). Although short-lived isotopic systems, most notably the lithophile–atmosphere $^{129}$I–$^{129}$Xe system ($t_{1/2} = 15.7$ Myr), the lithophile $^{146}$Sm–$^{144}$Nd system ($t_{1/2} = 103$ Myr), and the lithophile–siderophile $^{182}$Hf–$^{182}$W system ($t_{1/2} = 8.9$ Myr), were initially pursued primarily to date cosmochronological materials and processes, radiogenic decay of these systems has also led to the production of long-term isotopic heterogeneity in an array of sizable cosmochronological reservoirs, including the lunar and martian mantles (Nyquist et al., 1995; Foley et al., 2005; Debaille et al., 2009). Consequently, variations in the abundances of the daughter isotopes can also serve as tracers of planetary mantle mixing. Because the parent isotopes of these systems were extant for relatively short periods of time, ranging only from about 60 Myr for $^{182}$Hf, to as long as 600 Myr for $^{164}$Sm, the heterogeneities they record can only have formed early in solar system history. For example, the large differences in $^{182}$W/$^{184}$W and $^{142}$Nd/$^{144}$Nd isotopic ratios between nakhlite and shergottite meteorites, with both groups presumably derived from the terrestrial mantle (e.g., Caro et al., 2003, 2006; Willbold et al., 2011; Mukhopadhyay, 2012). Here, the term anomaly refers to an isotope composition that differs from that of the dominant composition for measurements of each isotope in terrestrial standards. Error bars associated with symbols reflect the 2σ analytical uncertainties for measurements of each isotope in terrestrial standards. 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for assessing the genetic makeup of materials added to planetary bodies during the final stages of assembly.

4. The final ~10 wt.% of accretion

The canonical model for the formation of the Moon involves a giant impact of an impactor comprising approximately 10% of the mass of the Earth (Canup and Asphaug, 2001). Although recent models for the putative giant impact have expanded the possible range for the mass of the impactor from as little as 5% (Ćuk and Stewart, 2012) to as much as 45% of the current mass of the Earth (Canup, 2012), there is little doubt that it was the final major accretionary addition to Earth. The event led to the transit of most of the metal from the impactor core through the Earth’s mantle. As a consequence, it could also have led to a substantial modification of siderophile element abundances and their isotopic compositions retained in the silicate portion of the Earth, following the impact (e.g., Halliday, 2004; Rubie et al., 2011). As evidenced by the mass balance constraints discussed above, subsequent late accretion could not have modified the abundances of the MSE in the mantle by more than ~10%. Thus, the giant impact would have been the last major event involved in the establishment of the modern characteristics of MSE in the terrestrial mantle. Consequently, some characteristics of MSE in the mantle today may provide insights to the nature of the final giant impactor involved in the construction of the Earth and Moon.

Of all the MSE, the modification of the mantle by giant impact has been most frequently studied using W isotopes, because of the radiogenic tracer capability inherent in the system (Halliday, 2004; Nimmo et al., 2010; Dwyer et al., 2014). Unfortunately, the W isotopic composition of the mantle today can presently provide only limited constraints regarding the nature of the giant impactor. This is because the outcomes of giant impact models involving the Hf–W isotopic system are strongly dependent on assumptions about the timing of the impact, and of core formation on both the proto-Earth and impactor. Of particular importance is the degree of equilibration that occurred between the core of the impactor and the silicate Earth, as metal from the impactor transited the mantle on its way to merge with the Earth’s core (e.g., Halliday, 2004, 2008). In contrast to the W present in metal derived from the core of the impactor, it is assumed that nearly all of the W present in the silicate portion of the impactor ended up mixing into the terrestrial mantle, as well as being added to the silicate portion of the Moon.

With regard to metal–silicate equilibration, one endmember possibility is that the core of the impactor quickly merged with the core of the Earth, with negligible exchange of W between the merging core and the silicate Earth (Fig. 4a). In this case, the giant impact would likely have led to an increase in the $\mu^{182}\text{W}$ (where $\mu^{182}\text{W}$ is the deviation in $^{182}\text{W}/^{184}\text{W}$ of a sample, compared to a modern terrestrial standard, in parts per million) of the Earth’s mantle as a result of the addition of W from only the silicate portion of the impactor (Halliday, 2004). This is based on the assumption that the $\mu^{182}\text{W}$ of the silicate portion of the impactor was more radiogenic than the proto-Earth’s mantle, because the embryo-size impactor is usually presumed to have accreted and differentiated a core and mantle more rapidly than Earth, creating a high Hf/W mantle, while $^{182}\text{Hf}$ was still plentiful (Dauphas and Pourmand, 2011). Thus, for this scenario, the $\mu^{182}\text{W}$ value of the silicate Earth could have risen 100 ppm or more as a result of the impact (Fig. 4a).

The other endmember possibility is that a substantial portion of the core of the impactor broke into small droplets, resulting from Rayleigh–Taylor instabilities, and the droplets equilibrated with the surrounding silicate liquid as they sank through the mantle to merge with the terrestrial core (Dahl and Stevenson, 2010). In this case, the W isotopic composition of the silicate Earth would likely have decreased significantly (Fig. 4b), as W with low $\mu^{182}\text{W}$ from the impactor core ($\mu^{182}\text{W} \leq -200$) equilibrated with the presumably more radiogenic silicate Earth (Halliday, 2008). In this case, the mantle of the Earth, just prior to the impact, may have been ≥200 ppm more radiogenic than the present mantle.

Because of these uncertainties in the extent of equilibration between metal and silicate, it is currently impossible to estimate the $\mu^{182}\text{W}$ value of the impactor mantle, as well as precisely determine the average age of core formation (where 50 wt.% of the core mass is achieved) for either the impactor or the proto-Earth. Nevertheless, W isotopes may have some utility in deconvolving the processes that occurred during and after the giant impact. Most importantly, it has been recently determined that the Moon has a $\mu^{182}\text{W}$ value that is enriched relative to the Earth’s mantle by 21–26 ppm (Kruijer et al., 2015; Touboul et al., 2015). The difference is most parsimoniously explained by assuming the W isotopic composition of the Earth and Moon were identical.

![Fig. 3. $^{100}\text{Ru}$ isotopic variations relative to terrestrial standards for a. iron meteorites and b. chondrites. The gray fields represent the 2σ analytical uncertainty for the measurements. Error bars associated with symbols reflect the 2σ analytical uncertainty for the measurements. The figure is modified from Chen et al. (2010).](image-url)
Immediately following the coalescence of the Moon, and that the small difference is the result of disproportional late accretion to the Earth and Moon, after the Moon formed. One major ramification of this observation is that, in order for the W isotopes and HSE abundances of the Earth and Moon to be self-consistent, the metal from the giant impactor equilibrated with the terrestrial mantle as the metal transited the silicate Earth (modified from Halliday, 2004). The increase in $\mu^{182}$W results from the addition of the radiogenic silicate portion of the impactor to the terrestrial mantle. b. The same model is used as for a, but for this scenario 100% of the core of the giant impactor equilibrates with the silicate Earth (modified from Halliday, 2008). The decrease in $\mu^{182}$W results from the addition of metal with appreciable W and very low $\mu^{182}$W (−200) from the core of the impactor to the terrestrial mantle.

This element. However, the uncertainties on the measured Mo isotopic compositions of enstatite chondrites have substantially different major element compositions (e.g., Mg/Si) compared to estimates for the composition of Earth's primitive mantle. So, unless there is a major hidden mantle reservoir with a substantially different bulk composition from the observed mantle, the enstatite chondrites cannot account for Earth's bulk composition (e.g., Palme and O'Neill, 2003).

For example, the $\Delta^{17}$O composition range of silicates present in IAB iron meteorites differs substantially from the Earth (Wasson and Kallemyn, 2002). Therefore, even if their Mo isotopic compositions overlap, the group is probably not closely genetically related to Earth.

Enstatite chondrites have been touted by some as potential building blocks of the bulk Earth (e.g., Javaux et al., 2010). Compositionally, however, it has been noted that the enstatite chondrites have substantially different major element compositions (e.g., Mg/Si) compared to estimates for the composition of Earth's primitive mantle. So, unless there is a major hidden mantle reservoir with a substantially different bulk composition from the observed mantle, the enstatite chondrites cannot account for Earth's bulk composition (e.g., Palme and O'Neill, 2003). That said, enstatite chondrites are good to excellent genetic matches to the isotopic composition of the bulk Earth with respect to, e.g., O, Ti, Ca, and Ni (Clayton et al., 1984; Carlson et al., 2007; Regelous et al., 2008; Trinquier et al., 2009; Herrwitz et al., 2014). This means that building blocks that were genetically similar to, but compositionally distinct from enstatite chondrites, are implicated in the construction of the bulk Earth. Not all isotopic compositions that reflect genetic componentry match, however. Fitoussi and Bourdon (2012) reported a −30 ppm difference in the Si isotopic composition between the silicate Earth and enstatite chondrites.

The Mo isotopic composition of enstatite chondrites overlap with the BSE within the present level of analytical precision (Burkhardt et al., 2011) (Fig. 2b). Burkhardt et al. (2011), however, noted that the characteristic w-shaped Mo isotope pattern for enstatite chondrites, as with ordinary and carbonaceous chondrites, is consistent with minor s-process depletion compared with the Earth. Thus, they concluded that the Mo isotopic compositions of enstatite chondrites are similar to, but distinct from, terrestrial Mo. Unlike the genetic similarities for elements like O, Ti, Ca and Ni, between the Earth and enstatite chondrites, which likely reflect the genetic history of the major building blocks of Earth, the Mo isotopic composition is most likely a reflection of the final 10–20 wt.% accretion. Thus, the fact that all of these elements have isotopic compositions similar or identical to enstatite chondrites suggests that >99% of the accretion of Earth occurred without a major change in the genetic signature of the materials contributing to growth.

Application of the MSe to study the nature of late stage building blocks is not limited to Earth. The Mo isotopic compositions of two spherulites analyzed by Burkhardt et al. (2011) overlap with the composition of the silicate Earth, so no genetic difference between these two bodies can be detected at the current level of analytical resolution for this element. However, the uncertainties on the measured Mo isotopic
composition for these samples are relatively large. Future refinement of the Mo isotopic composition of Mars, and determination of whether or not it has a uniform isotopic composition, will be critical to assessing whether late stage additions to the Earth and Mars came from genetically similar materials, and whether late stage building blocks to Mars were well homogenized before the large-scale differentiation events that led to isotopic heterogeneity in $^{182}\text{W}$ and $^{142}\text{Nd}$.

In summary, the isotopic compositions of W in terrestrial and lunar rocks currently provide little direct information about the giant impactor that was the final major building block of the Earth. The data do, however, provide important new constraints on the physical processes that occurred during this final major accretionary event. Most importantly, it appears that metal from the impactor efficiently stripped HSE from the proto-Earth's mantle, such that the late accretionary clocks for the reconstituted Earth, and newly coalesced Moon, began at the same time. The Mo isotopic composition of the terrestrial mantle is likely dominated by Mo provided by the building blocks that contributed the final 10–20 wt.% to Earth's mass. The Mo isotopic composition of the mantle is very similar to enstatite chondrites, much like isotopic compositions of other genetic tracers that reflect the composition of the bulk Earth. This suggests that the building blocks of Earth were genetically similar to enstatite chondrites, and did not vary much from a genetic standpoint over the course of the first >99% of Earth's construction.

5. The final ~0.5 wt.% of accretion

5.1. Highly siderophile elements in the bulk silicate Earth

The absolute and relative abundances of HSE in primitive meteorites vary among the major chondrite groups (Walker et al., 2002a; Horan et al., 2003; Brandon et al., 2005; Tagle and Berlin, 2008; Fischer-Gödde et al., 2010) (Fig. 5a–d). For example, Pd/Ir in some enstatite chondrites tend to be higher than for ordinary, carbonaceous, or R-type chondrites. Given that Re/Os also varies among the chondritic groups, the $^{187}\text{Os}/^{188}\text{Os}$ of chondrites can serve as an important, complementary parameter to discriminate among chondrite groups. Most notably, carbonaceous chondrites are characterized by generally lower $^{187}\text{Os}/^{188}\text{Os}$, and therefore lower long-term Re/Os, than ordinary and enstatite chondrites (Fig. 6). Modest, long-term differences in Pt/Os have also resulted in small differences in $^{180}\text{Os}/^{188}\text{Os}$ among some chondrite groups (Brandon et al., 2006), although the current level of resolution is insufficient for this system to be very useful as a tracer.

Although variations in the relative abundances of HSE are becoming increasingly better constrained for chondritic components (e.g., Horan et al., 2009; Archer et al., 2014), except for calcium–aluminum rich inclusions, there has been only limited success in determining the causes of the variations among the HSE characteristics of bulk chondrites (Mason and Taylor, 1982; Sylvester et al., 1990; Fischer-Gödde et al., 2010). Thus, no nebular or parent body processes can currently be firmly linked to the HSE characteristics observed in bulk chondrites. This means the process that led to lower, long-term Re/Os in carbonaceous chondrites, compared to other chondrite groups, may not be related to the accompanying, generally more volatile-rich nature of carbonaceous chondrites. Despite these current limitations to fingerprinting late accretionary additions, if most of the mass of HSE present in the BSE today was added as a result of late accretion, the relative abundances of these elements in the BSE should provide an averaged compositional snapshot of the final ~0.5% of mass addition to Earth.

Establishing HSE abundances in the BSE with sufficient precision to make comparisons to possible cosmochemical precursors is problematic.

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Fig. 5. a–d. Variance in HSE ratios and $^{187}\text{Os}/^{188}\text{Os}$, which serves as a proxy for long-term Re/Os, in the major primitive meteorite groups. Data for bulk chondrites are from Walker et al. (2002a), Horan et al. (2003), Brandon et al. (2005) and Fischer-Gödde et al. (2010). Also shown is the estimate for the BSE from Becker et al. (2006).
Because of the sensitivity of $^{187}\text{Os}/^{188}\text{Os}$ ratios to discriminate among small, long-term differences in Re/Os, Os isotopes have been especially heavily used among the HSE to characterize the Re/Os of the mantle (Hirt et al., 1963), and ultimately, the nature of late accreted materials to Earth (Morgan, 1985; Meisel et al., 1996, 2001). However, the application is not straightforward. Although Os is highly compatible within the mantle, the Os isotopic composition of the BSE cannot be measured directly via analysis of the dominant silicate reservoir in Earth, the oceanic mantle. This is because Re behaves incompatibly during mantle melting (Barnes et al., 1985; Rehkämper et al., 1999; Pearson et al., 2004). Consequently, oceanic and continental crustal extraction has modified the Re/Os of the residual oceanic mantle over Earth history. Abundances of Re estimated for the continental crust are relatively low (Peucker-Ehrenbrink and Jahn, 2001), and given the limited mass of this reservoir, its formation is unlikely to have led to significant modification of the Re/Os of the residual mantle. By contrast, the formation of oceanic crust, with its comparatively high Re concentrations, may have significantly modified the Re/Os of the residual mantle. The magnitude of this modification is open for debate, as some of the Re extracted into oceanic crust has been recycled back into the mantle. How much of the Re (and Pt) has been re-mixed back into the oceanic mantle remains difficult to assess (Walker et al., 2002b).

To circumvent this problem, Meisel et al. (1996, 2001) applied a projection method utilizing the compositions of variably melt-depleted mantle peridotite xenoliths to estimate the $^{187}\text{Os}/^{188}\text{Os}$ of the BSE. They plotted $^{187}\text{Os}/^{188}\text{Os}$ (record of long-term Re/Os) versus Al$_2$O$_3$ or Lu, indicators of mantle melt depletion that are not as easily modified by secondary processes as Re, and projected the resulting linear trends to points of intersection with assumed Al$_2$O$_3$ or Lu concentrations in the BSE. Using this method, Meisel et al. (2001) reported a $^{187}\text{Os}/^{188}\text{Os}$ ratio of 0.1296 ± 0.0008 (2σ). This ratio must be considered a minimum estimate because the projections were made mainly from samples derived from sub-continental lithospheric mantle. Such peridotites must have been physically separated from the oceanic mantle at some time prior to isolation as sub-continental lithospheric mantle. The oceanic mantle itself was most likely variably depleted in Re prior to these reservoirs transitioning from oceanic to sub-continental lithospheric mantle, so the melt depletion events recorded in these rocks likely included at least one stage of prior melt depletion. The $^{187}\text{Os}/^{188}\text{Os}$ estimated for the BSE is at the upper end of the range of compositions recorded in bulk ordinary and enstatite chondrites (Fig. 6). Given this, the $^{187}\text{Os}/^{188}\text{Os}$ of the BSE appears to be most similar to ordinary and enstatite chondrites, or even slightly more radiogenic, and this likely means that carbonaceous chondrites, or similar materials, were not major players in late accretion (Walker et al., 2002a). This conclusion has subsequently been supported by arguments based on isotopic evidence for other elements (e.g., Jacobson et al., 2014). Given the volatile-rich nature of some carbonaceous chondrite groups, this observation in turn has been taken as evidence that late accretion provided little water to Earth (Drake and Righter, 2002), although as noted, there is currently no known process that relates the incorporation of volatiles and low Re/Os into the parent bodies of carbonaceous chondrites.

Absolute and relative abundances of other HSE have been estimated for the BSE using a similar approach as for $^{187}\text{Os}/^{188}\text{Os}$ (Becker et al., 2006; Fischer-Gödde et al., 2011). When collectively considering the HSE characteristics estimated for the BSE as compared with chondrites (Fig. 7), it appears to be most similar to enstatite chondrites for Re/Os, Os/Ir, Pt/Ir and Pd/Ir (Becker et al., 2006). However, the BSE appears to be modestly suprachondritic with respect to Ru/Ir and possibly Pd/Ir (Becker et al., 2006). It is possible that the slight Pd enrichment could be the remnant effect of high pressure–temperature metal–silicate partitioning as discussed with regard to the MSE (e.g., Mann et al., 2012). Current experimental data, however, provide no evidence that this could also be true for Ru. Thus, although the current estimate of the HSE composition of the BSE is most like enstatite chondrites, it is not a perfect match to any known chondrite group, or individual chondrites present in our collections. This could mean that late accretion to Earth was dominated by one or more components that were somehow processed in the nebula, or on a parent body, such that HSE were fractionated relative to presently sampled chondrites. This in turn could occur as a result of the location of formation within a chemically heterogeneous protoplanetary disk, or formation at a different time in a chemically evolving nebula. Given that there are bulk chondrites with individual HSE ratios that extend beyond those estimated for the BSE (although no one chondrite has all of the HSE characteristics of the BSE), nebular or parent body processes can evidently cause such fractionations. It is also possible that the fractionated HSE abundances present in the dominant late accretionary component could result from another process, such as crystal–liquid fractionation of a metal component (e.g., Fischer-Gödde et al., 2011).

It is important to recognize that there are some limitations to the projection approach towards characterizing the BSE and fingerprinting late accretionary additions, using HSE. Of greatest concern is the precision and accuracy of HSE estimates for the BSE. The absolute and relative concentration estimates in the BSE are based on measurements of mantle peripheralities that may represent the end stage of multiple processes, including mantle melting, metasomatism and crustal recycling (Alard et al., 2000; Le Roux et al., 2007). Thus, although the broadly chondritic nature of the BSE is of little doubt, the relatively small differences in elemental ratios that are key to discriminating among chondritic groups, or fingerprinting a heretofore unidentified chondrite-like contributor to
the mantle, have been called into question (e.g., Lorand et al., 2009). The primary question is whether any mantle peridotites can provide sufficient fidelity in recording the BSE composition. The typically strongly compatible natures of Os, Ir, and Ru, make them less susceptible to modification by partial melting and metasomatic processes, compared to Pt, and especially the incompatible Pd, Au, and Re. Consequently, they provide the strongest constraints on the HSE composition of the BSE, and the suprachondritic nature of Ru/Ir in the BSE appears to be real. Further, the similarity of HSE characteristics of large numbers of variable “fertile” mantle peridotites, ranging from abyssal peridotites, to peridotites from the mantle sections of ophiolites, to oceanic mantle xenoliths, supports the contention that the BSE is characterized by suprachondritic Ru/Ir and possibly Pt/Ir (Becker et al., 2006; Fischer-Gödde et al., 2011). However, compilation of an even larger number of data from all types of mantle lithologies, combined with an improved understanding of how HSE from oceanic crust have been recycled back into the oceanic mantle, may ultimately be required to assemble a high-confidence understanding of HSE in the BSE.

The most promising element for examining the genetic characteristics of late accreted HSE is Ru. As a HSE, Ru was strongly concentrated into metal by core formation processes. As noted above, the large range of nucleosynthetic isotopic compositions recorded in meteorites (Chen et al., 2010) makes this element ideal as a genetic tracer of cosmic materials contributed by late accretion. To date, Ru data for terrestrial rocks and meteorites are very limited. However, the most recent results (Fischer-Gödde et al., 2014; Bermingham et al., 2015) suggest that, as with Mo, the isotopic composition of Ru in the BSE is very similar to enstatite chondrites, and is well resolved from all carbonaceous chondrites. This observation is then consistent with the Os isotopic results.

5.2. Constraining the physical nature of late accretion

One key aspect of utilizing siderophile elements as genetic tracers of planetary building blocks requires knowledge of the specific physical processes involved in their incorporation into planetary mantles. This is especially true for the HSE. For example, late accretion of HSE to Earth’s mantle may have occurred as a consequence of a relatively gentle rain of smaller bodies onto the surface of the planet, as envisioned by some workers (Anders, 1968; Turekian and Clark, 1969). This concept led to the coinage of the term late veneer, as discussed above. In this case, a dog’s breakfast of HSE-rich materials of diverse geneticities may have been slowly mixed downward into the mantle as a result of crustal recycling, coupled with mantle convection. One way to test this hypothesis would be to look at the evolution of HSE abundances in the Archean mantle using high-degree partial melts that extract large proportions of HSE from the mantle and experience little to no fractionation en route to the surface, such as komatiites. Komatiites are high-MgO lavas, which are commonly presumed to be derived from partial melting in deep mantle upwellings, or plumes (Campbell et al., 1989). Maier et al. (2009) argued that gradual downward mixing of late accreted materials can potentially account for the low abundances of HSE in some early Archean komatiites, and that the “normal” HSE abundances found in komatiites formed later than ~2.9 Ga indicate that the putative late accreted materials had become well-mixed into the deep mantle sources of komatiites by that time.

The combined HSE abundances and Re–Os and Pt–Os isotopic studies have resulted in robust estimates of HSE abundances in deep mantle sources of komatiites, ranging in age from 3.6 to 2.4 Ga using the projection technique (Puchtel et al., 2004a, 2004b), whereas the $^{187}$Re–$^{187}$Os and $^{198}$Pt–$^{186}$Os isotopic systematics confirmed post-crystallization, closed-system behavior of the rocks with respect to HSE abundances (Puchtel et al., 2009a, 2009b). The Os isotopic evidence for closed system behavior provides some additional confidence that the projected mantles source abundances are accurate. The relative HSE abundances obtained for the late Archean komatiite systems were concluded to be best matched by those in enstatite chondrites (e.g., Puchtel et al., 2004a, 2009a). The komatiite systems studied in this manner show significant variations in the absolute HSE abundances between the sources of late Archean komatiite systems (Fig. 8), but are generally similar to those in the sources of early Archean komatiite systems, indicating little temporal changes in the HSE abundances (Puchtel et al., 2014). There are notable exceptions, however, with the oldest early Archean komatiite system (3.55 Ga Schapenburg) having substantially lower HSE abundances compared to all other komatiite systems studied to-date, which was interpreted to reflect sluggish mixing of post-magma ocean domains in the Archean mantle (Puchtel et al., 2009b).

Another way to advance understanding of the physical nature of late accretion is by combining observations of the chemical characteristics of planetary materials with dynamical models for the first ~500 Ma of solar system history. Here, comparing the characteristics of the HSE present in the terrestrial mantle with abundances present in the lunar and martian mantles may be particularly important. The abundances and isotopic compositions of the HSE in at least the upper portion of the terrestrial mantle are generally well-defined and limited in variation. As noted, the $^{187}$Os/$^{186}$Os and $^{186}$Os/$^{188}$Os ratios estimated for the BSE are within the range of chondritic meteorites (Walker et al., 1997: Meisel et al., 2001; Brandon et al., 2006). Further, data for the global sampling of oceanic and subcontinental lithospheric mantle peridotites are characterized by relatively uniform abundances of Os and Ir, two HSE that are highly compatible during partial melting of the mantle (e.g., Rehkämper et al., 1999; Morgan et al., 2001). While there has long been a question as to whether the upper mantle is more enriched in HSE than the lower mantle, seismic tomography over the past 20 years has documented the exchange of matter between upper and lower mantle, providing evidence for mantle plumes rising from the lower mantle and subducting slabs transiting into the lower mantle (e.g., Goes et al., 1999; Nolet et al., 2006). It is, therefore, likely that there are presently no major, global-scale HSE concentration variations within the mantle. Consequently, uncertainty in the mass of late accreted materials to the mantle, necessary to account for the observed abundances of HSE, primarily reflects the factor of two variation in HSE.

Fig. 8. Calculated total Pt + Pd abundances in the sources of Archean komatiites as compared to the BSE estimate of Becker et al. (2006). For each of these systems, $^{187}$Re–$^{187}$Os isochrons consistent with the known ages of the rocks have been generated, consistent with closed-system behavior for other HSE. The late Archean komatiite systems span a substantial range, from 58 ± 7% in the 2.69 Ga Belingwe system to 85 ± 5% in the 2.72 Ga Abitibi system, of the total Pt and Pd present in the estimates for the modern BSE, with the 2.41 Ga Vetryen system being at the lower end of this range. Within the uncertainties, the total Pt + Pd abundances in some of the late Archean komatiite systems, e.g., Abitibi and Kostomuksha, overlap with those in the estimates for the BSE, whereas in the others (Vetryen, Belingwe, and Volotok–Kamennoozero) fall slightly (by 20–14%) if the full uncertainties on the estimates are considered) short of reaching that level. The total Pt + Pd abundances in the sources of the two of three early Archean komatiite systems from the Barberton Greenstone Belts, South Africa, are within the range of those in the late Archean komatiite systems, albeit on the lower side, varying from 56 ± 12% for the Komati to 65 ± 10% for the Weltevreden, of the total Pt + Pd present in the estimates for the modern BSE, whereas the third system (Schapenburg) is characterized by much lower Pt + Pd abundances (27 ± 4% of BSE). Data sources: Puchtel et al. (2004a,b, 2005, 2007, 2009a,b, 2014), and Puchtel and Humayun (2005).
abundances between different types of chondritic meteorites (e.g., Horan et al., 2003), rather than uncertainties in the HSE content of the mantle. It is much more difficult to constrain the abundances of the HSE in the lunar and martian mantles compared to the terrestrial mantle. No mantle samples have, as yet, been collected from either body. Hence, mantle abundances of the HSE must be deduced from mantle-derived volcanic rocks. Based on HSE abundances present in leached samples of picritic glass spherules and lunar basalts, Walker et al. (2004) and Day et al. (2007), respectively, estimated that HSE concentrations in the lunar mantle are a factor of 20 or more lower than in the terrestrial mantle. Day et al. (2007), respectively, estimated that HSE concentrations in the lunar mantle are a factor of 20 or more lower than in the terrestrial mantle (Fig. 9a). Such a large difference in concentration, if correct, cannot be explained by gravitational focusing or inefficiency of impactor retention. The lower concentrations may instead reflect proportionally much less mass added to the lunar mantle by late accretion, compared to the Earth. This in turn could result from a longer period of late accretion for Earth, compared to the Moon, i.e., one that began well before formation of the Moon. However, as noted above, recent W isotopic data for the Moon suggest that the Moon-forming giant impact efficiently removed HSE from the mantle to Earth’s core and reset the late accretionary clock for the two bodies (Kruijer et al., 2015; Touboul et al., 2015). If so, this means that the dominant late accretionary periods of the Earth and Moon began after formation of the Moon, and were contemporaneous. Unless substantial quantities of late accreted materials to the Moon are sequestered from our view in its lower crust (e.g., Schlichting et al., 2012), an interpretation for which there is currently no physical evidence, there remains a sizable mismatch in the proportions of late accreted materials added to the mantles of the Earth and Moon. As noted by Walker (2009) this mismatch presents a potential problem for the late accretion hypothesis.

In contrast to the lunar mantle, the martian mantle appears to have HSE abundances that are surprisingly similar to those present in the terrestrial mantle. Basaltic and ultramafic shergottite meteorites, commonly believed to come from Mars, are characterized by HSE abundances that scale with the MgO content of the rocks in a way that is similar to terrestrial volcanic rocks (Brandon et al., 2012) (Fig. 9b). In addition, shergottites exhibit a range of initial $^{182}\text{W} / ^{184}\text{W}$ that is very similar to the range of compositions present in terrestrial mantle-derived volcanic rocks (Brandon et al., 2012). Given the fact that Mars is commonly presumed to have formed as a planetary embryo prior to the Earth (Dauphas and Pourmand, 2011), it might be expected to have a larger proportion of late accreted material mixed throughout its mantle. Instead, all existing data suggest that the HSE concentrations in the martian mantle are similar to those in the terrestrial mantle, indicating that a roughly similar proportion of late accreted materials was added to the mantle of Mars (Brandon et al., 2012).

Bottke et al. (2010) reported that one way to account for the similarity of HSE abundances in the terrestrial and martian mantles, but much lower HSE abundances in the lunar mantle is by a process they termed stochastic late accretion. The principle of stochastic late accretion is based on the assumption that most late accretionary mass was added to the Earth and Mars by a very limited number of impacts of approximately Pluto mass bodies (~$1 \times 10^{22}$ kg). By chance, the Moon was not struck by any bodies of this size, and so retained relatively low abundances of HSE. Subsequent dynamical models have highlighted the probability that bodies of similar mass may have survived beyond the formation age of the Moon, and thus been available to participate in stochastic late accretion (Marchi et al., 2014). Bottke et al. (2010) reported that for impactors with diameters between 2000 and 2500 km, slightly more than 10% of their dynamical simulations could achieve the necessary relative enrichments in HSE to the mantles of the Earth, Moon and Mars.

If stochastic late accretion correctly accounts for the apparent disparity in HSE abundances between the lunar and terrestrial mantles, it has a major implication for tracing late stage building blocks of the Earth, and possibly Mars. It requires that mass was added to the Earth by a limited number of impact events that likely would have generated discrete magma seas or lakes, rather than as a chemically and isotopically well-mixed veneer of small bodies. Consequently, if late stage impactors were added to the mantle in such a way that global melting did not occur, then the impactors may have imparted isotopically distinct HSE signatures to different portions of the mantle, assuming that the impactors were genetically different from the average BSE, and from one another.

But what is the likelihood that moderately-sized, early-formed mantle heterogeneities remained isotopically distinct for hundreds of millions of years, until they melted to produce rocks that became incorporated into the rock record? This is where it is important to consider data for the short-lived radiogenic isotope systems. Perhaps of greatest importance for consideration here are $\mu ^{182}\text{W}$ isotopic data for rocks that were ultimately derived from the terrestrial mantle. Anomalous $\mu ^{182}\text{W}$ values have been identified in a number of ancient rocks, including $\geq 3.8$ Ga supracrustal rocks from Nuvvuagittuq, Quebec (Touboul et al., 2014), ~3.7 Ga supracrustal rocks from Isua, Greenland (Willbold et al., 2011), and 2.8 Ga komatiites from Kostomuksha, Fennoscandia (Touboul et al., 2012). All $^{182}\text{W}$ anomalies for terrestrial rocks, reported to date, range between +5 and +15 ppm (Fig. 10). These terrestrial anomalies are much smaller than the isotopic variations that have been observed for other early solar system objects, which range from initial $\mu ^{182}\text{W}$ values determined for calcium aluminum-rich inclusions of ~350 ppm, to enriched values of as much as +4000 ppm in some eucrites (Kleine et al., 2009).
Terrestrial enrichments in $^{182}\text{W}$ have been interpreted in two different ways. Willbold et al. (2011) reported $^{182}\text{W}$ enrichments averaging ~13 ppm for 3.7 Ga supracrustal rocks from Isua, Greenland. These authors proposed that the enriched compositions reflect derivation of precursor rocks from a mantle domain that formed prior to a final major stage of late accretion, and that this mantle domain remained mostly free of late accreted materials until it melted to form the Isua rocks. Thus, the mantle precursor materials to the Isua rocks formed by normal planetary accretion, were stripped of HSE by metal segregation during core formation, then remained isolated from HSE and W (with low $\mu^{182}\text{W}$) added by subsequent late accretion. This is a process that could lead to isotopic heterogeneity in the mantle long after $^{182}\text{Hf}$ became extinct. Most importantly, this mantle domain was not contaminated with late accreted HSE as a result of mantle mixing until after the early Archean melting event that produced the Isua precursor rocks, presumably well after completion of the dominant phase of late accretion. This requires inefficient mixing of the mantle during the Hadean through early Archean. If this interpretation is correct for the Isua rocks, then the mantle domain sampled by them should have been relatively devoid of HSE. Willbold et al. (2011), however, did not report complementary HSE for these rocks. Even when this is eventually done, however, it should be recognized that constraining the concentrations of HSE in the mantle source(s) of such highly altered supracrustal rocks will be challenging.

A second means to account for anomalous $^{182}\text{W}$ in mantle-derived rocks is by solid–liquid fractionation processes that may have occurred in the mantle while $^{182}\text{Hf}$ was still extant. Because the absolute HSE abundances estimated for the mantle source of the Kostomuksha komatiites are nearly identical, within uncertainties (Puchtel and Humayun, 2005), to those in the BSE estimates of Becker et al. (2006), Touboul et al. (2012) rejected the model of Willbold et al. (2011) for these rocks. They instead concluded that either metal–silicate fractionation in a basal magma ocean, or silicate crystal–liquid fractionation in a more conventional, whole mantle magma ocean led to the creation of a mantle domain characterized by high Hf/W. This in turn led to the formation of a domain that evolved to high $\mu^{182}\text{W}$, as $^{182}\text{Hf}$ decayed. Because of the short lifetime of $^{182}\text{Hf}$, it was concluded that the fractionation events occurred within the first 30 Myr of solar system history. Similar processes may also have led to the creation of some $^{142}\text{Nd}$ anomalies (e.g., Brown et al., 2014).

Regardless of the true mechanisms involved in the generation of terrestrial $^{182}\text{W}$ anomalies, it is clear that their presence in the rock record requires the long term survival of chemical heterogeneities in the terrestrial mantle, despite presumably vigorous convective mixing. Much larger $^{182}\text{W}$ and $^{142}\text{Nd}$ isotopic anomalies have been determined for some, but not all martian meteorites, so the martian mantle also likely escaped a final large-scale homogenization event during the final stages of its growth. Thus, if these bodies experienced late stages of accretion from genetically diverse materials, it might be expected that attenuated signals from the various materials might be summoned from the rock record. Limited high precision analyses of Ru isotopes in terrestrial materials have, as yet, not identified any isotopic

![Fig. 10. Plot of $\mu^{182}\text{W}$ values for ancient rocks from the ≥3.8 Ga Nuvvuagittuq, Quebec, supracrustal suite, the ~3.7 Ga supracrustal rocks from Isua, Greenland, the 3.5 Ga Komati komatiites from the Barberton Greenstone Belt, South Africa, and 2.8 Ga komatiites from Kostomuksha, Fennoscandia. Data are from Willbold et al. (2011), Touboul et al. (2012) and Touboul et al. (2014).](image)
heterogeneity within the mantle (Bermingham et al., 2015), but the search has just begun.

5.3. The Mo–Ru connection

In the discussion above, it is noted that the Mo and Ru present in the mantle today were most likely emplaced by different late-stage accretionary processes. The Mo abundance of the mantle was dominantly established by the final 10–20% of terrestrial accretion, with major input from the giant impact stage of growth, culminating in the Moon-forming giant impact. The Mo isotopic composition of the mantle probably represents a mixture of Mo from the silicate portion of the proto-Earth, as well as Mo from both the core and mantle of the giant impactor. By contrast, most Ru was likely added to the mantle by late accretion. Dauphas et al. (2004, 2014) made the important observation that, when plotting the range of nucleosynthetic heterogeneities for Mo and Ru connection

Fig. 11. Plot of $^{260}$Mo versus $^{190}$Ru of group averages chondrites and iron meteorites showing the linear trend defining s-process enriched and depleted compositions. Note that Earth plots at one end of the trend. Data are from Burkhardt et al. (2011) and Chen et al. (2010).

accretion, whereby late accretionary mass to the Earth and Mars was primarily added through the impact of several $10^{22}$ kg mass objects, whereas the Moon was not struck by such large impactors.

The HSE characteristics of the BSE, as assessed by the relative abundances of the HSE (including $^{187}$Os/$^{188}$Os) are most like enstatite and ordinary chondrites, although the HSE characteristics of the BSE are not a perfect match to any chondrite group. Osmium isotopes strongly suggest that the late accreted materials to Earth were not like carbonaceous chondrites. Although still very limited for both terrestrial rocks and meteorites, Ru isotopic composition data are also consistent with a dominant late accretionary component to the Earth that is genetically similar to enstatite chondrites. Remarkably, this may indicate that the genetic make-up of even late-accreted materials to Earth, did not diverge substantially from the basic building blocks of the Earth.

6. Late heavy bombardment: the final $\sim$0.05 wt.% of accretion?

It has long been hypothesized that the Earth–Moon system, and likely the entire inner solar system, underwent a phase of late accretion, termed late heavy bombardment (LHB), within the interval from $\sim$4.1 to $\sim$3.8 Ga. The evidence for this putative event primarily comes from geochronologic information obtained from a variety of shocked and/or melted lunar rocks (e.g., Turner et al., 1973; Tera et al., 1974; Kring and Cohen, 2002). For example, Tera et al. (1974) recognized that most rocks collected by the Apollo missions formed a linear trend on a plot of $^{207}$Pb/$^{206}$Pb versus $^{238}$U/$^{204}$Pb that intersects concordia at about 3.9 Ga. Because of the ubiquity of this age, they inferred that the Moon underwent what they referred to as a terminal cataclysm. They envisioned the cataclysm to have been a relatively brief period of heavy bombardment ($<$300 Ma) during which the surface of the Moon, and presumably the Earth, was peppered with large impactors, leading to the creation of at least some of the lunar basins. Current estimates suggest the basin-forming impactors were as large as 200 km in diameter (e.g., Hurwitz and Kring, 2014). Subsequent studies of lunar impact melt rocks have provided strong support for a major disturbance in ages at about 3.9 Ga (e.g., Dalrymple and Ryder, 1993; Cohen et al., 2000), as few impact-modified lunar rocks yield ages older than $\sim$3.9 Ga. Although the LHB had a major effect on shaping the surface of the Moon, it likely involved much less mass than is envisioned for late accretion as a whole. Even generous estimates for the mass of the LHB, based on assumptions about the masses of basin-forming impactors, place the mass of materials involved at no more than about 10% of estimates for the overall mass of late accretionary additions (0.05 wt.% accretion) (e.g., Morgan et al., 2001), and some dynamical models suggest that only about 1% (0.005 wt.% accretion) was added by the LHB (Morbidelli et al., 2012; Marchi et al, 2014).

Dynamical models for the evolution of the solar system have suggested some possible causes for a period of LHB (Morbidelli et al., 2001; Gomes et al., 2005; Strom et al., 2005; Chambers, 2007). For example, Gomes et al. (2005) suggested that migration of Uranus and Neptune, resulting from Jupiter and Saturn entering a 2 to 1 orbital resonance, may have led to the perturbation in the orbits of small bodies from both the asteroid belt and the Kuiper belt. Chambers (2007) suggested that the removal of a now missing additional planet, between the orbit of Mars and the asteroid belt, could have triggered the LHB. Despite these observations and models, it is also possible that the – 3.9 Ga age represents a re-set age for the samples from a limited areal extent on the near-side of the Moon, from which all Apollo samples were collected, or even sampling of ejecta from only the youngest of the major basins (Spudis et al., 2011).

In addition to seeking to understand the timing of the LHB, it is also imperative to constrain the chemical characteristics of the materials involved in the bombardment because of the possibility that they delivered substantial water and other volatile species to the Earth and Moon. The primary means to examine the chemical characteristics of materials from the putative LHB has been to analyze lunar impact melt rocks that
were created as a result of the basin-forming impacts. Such studies have been pursued since the Apollo missions, with chemical characterizations focused upon siderophile elements (e.g., Morgan et al., 1972, 1974; Korotev, 1994). Siderophile element data for nearly all studies prior to ca. 2000 were obtained by neutron activation analysis, so elements such as Ir, Au, Ni, and Ge, that can be well-measured by this method, were most commonly considered. For example, Morgan et al. (1974) concluded that rocks from the Apollo 17 site included meteoritic components from at least six impactors, none of which had siderophile element characteristics that perfectly matched known meteorites.

Such studies as Morgan et al. (1974) assumed that endogenous lunar highlands or basaltic rocks formed with very low siderophile element abundances, so that the siderophile elements present in the impact melt rocks were nearly entirely derived from one or more large impactors. A substantial number of studies reporting siderophile element data for so-called pristine lunar rocks have generally borne out this assumption (e.g., Warren and Wasson, 1977; Ryder et al., 1980; Warren et al., 1991; Day et al., 2007, 2010).

Although the early studies provided highly valuable insights into the chemical nature of impactors, some of the chief elements used to discriminate among possible impactors, such as Au and Ge, are moderately volatile and could potentially have been modified by high temperature impact processes. To circumvent this problem, Norman et al. (2002) first applied the isotope dilution technique, teamed with inductively-coupled plasma mass spectrometry, to measure a larger suite of HSE in Apollo 17 impact melt rocks. That study measured and reported data for Re, Ir, Ru, Pt, and Pd, and identified at least three sources of HSE to the Apollo 17 suite. One source had HSE ratios similar to ordinary chondrites. A second component was characterized by HSE similar to EH chondrites. In order to account for suprachondritic Re/Ir, Ru/Ir, and Pd/Ir in most of the rocks, Norman et al. (2002) appealed to the possibility of a third component, either an endogenous component enriched in Re, Ru, and Pd, or an older component in the target crust that was incorporated into the crust by an earlier impactor with non-chondritic relative abundances of HSE.

Four recent, subsequent studies have utilized similar isotope dilution techniques to measure the abundances of Re, Os, Ir, Ru, Pt, and Pd in lunar impact melt rocks, as well as measure $^{187}$Os/$^{188}$Os, which serves as a sensitive proxy for long-term Re/Os (Puchtel et al., 2008; Fischer-Gödde and Becker, 2012, 2014; Liu et al., 2015). A major difference between these studies and the study of Norman et al. (2002) is that they examined multiple pieces of each rock studied. In approximately half of the rocks examined, the resulting plots of Ir versus each of the other HSE measured yielded linear trends with intercepts indistinguishable from 0, within regression uncertainties (Fig. 12). In such cases, the trends can be assumed to represent mixing between the exogenous impactor and the HSE poor lunar target rocks, similar to interpretations for terrestrial impact melt rocks (e.g., McDonald et al., 2001). The slopes of the linear trends can, therefore, be assumed to record the HSE ratios of the basin forming impactors.

Puchtel et al. (2008) and Sharp et al. (2014) reported and interpreted data mainly for Apollo 17 impact melt rocks. Both studies reported a “dominant” component for the site, most notably characterized by suprachondritic Re/Os (as measured by $^{187}$Os/$^{188}$Os), as well as Ru/Ir and Pd/Ir comparable to the results from Norman et al. (2002). They interpreted the results to suggest that the dominant source of HSE to the site, most likely the spatially associated Serenitatis basin impactor, shared broad similarities to some chondritic meteorites (enstatite chondrites), but sampling a composition not presently found in our meteorite collections. By contrast, a feldspar rich, or granulitic component present as clasts in some of these rocks, was determined to be characterized by relative abundances of HSE more similar to carbonaceous and ordinary chondrites. Fischer-Gödde and Becker (2012) focused mainly on impact melt rocks from the Apollo 16 site. Here, they found Re/Os, Ru/Ir, Pt/Ir, and Pd/Ir ratios extending much higher than in known chondrites, and even well beyond the range of Apollo 17 rocks. They also analyzed some granulitic rocks and reported that, like prior studies, this component is most like ordinary chondrites. Of note, this study recognized that virtually all of the HSE data for Apollo samples plot along linear trends of HSE/Ir versus $^{187}$Os/$^{188}$Os. They interpreted this to mean that all of the Apollo impact melt rocks incorporated at least two HSE-rich components at the time of their formation. One was very similar to carbonaceous chondrites and is the major component in granulitic rocks. The other component resembles a chemically evolved group IVA iron meteorite. Consequently, they proposed that both components became variably mixed during basin-forming impacts, but were not substantially modified by the inclusion of HSE derived from the basin-forming impactors.

Most recently, data from Liu et al. (2015) for Apollo 15 and 16 melt rocks filled in the gaps in the apparent linear trend recognized by Fischer-Gödde and Becker (2012), thus, strengthening their observation. In the compilation of data reported by Liu et al. (2015), nearly all data for lunar impact melt rocks plot along a continuous linear trend ranging from a HSE composition that is broadly chondritic, to an endmember with $^{187}$Os/$^{188}$Os, Ru/Ir and Pd/Ir ratios far above those of known chondrites (Fig. 13a-d). Two possible scenarios to explain the observed trends are: 1) Variable mixing between an earlier granulitic contaminant and a series of later-stage impactors that happened to form co-linear, suprachondritic Re/Os, Ru/Ir, Pt/Ir and Pd/Ir; 2) Variable mixing between two components present in the lunar crust prior to the late-stage basin forming impacts. For this scenario, one component was chondritic in composition and the other component had fractionated HSE, and could have been a core fragment, as suggested by Fischer-Gödde and Becker (2012). Although the latter scenario requires the involvement of a portion of an evolved core, it currently appears to be the most simplistic explanation for the trend. It also requires that the later-stage basin forming impacts (e.g., Imbrium) added only very limited HSE to the sampled impact melt rocks from multiple sites. These models await genetic testing using the nucleosynthetic anomalies characteristic of siderophile elements Mo and Ru.

In summary, the HSE characteristics of lunar impact melt rocks, believed to have formed as a result of basin forming events during a possible LHB, include at least two major components. One, with HSE characteristics similar to carbonaceous or ordinary chondrites, the other component is much more highly fractionated and most resembles a portion of a core represented by some iron meteorites. One way to
account for the data is to conclude that both components were incorporated in the lunar crust prior to the LHB, and that the basin-forming impacts mainly resulted in variable mixing of the older components. No isotopic data, e.g., Ru, are as yet available for genetic testing of these components.

7. Putting it all together

At the present time, combined lithophile and siderophile element data suggest that the primary building blocks of Earth were isotopically broadly similar to enstatite chondrites, but probably substantially differed from enstatite chondrites in terms of major element composition. Combining isotopic data for lithophile and siderophile elements that serve as genetic tracers leads to the suggestion that there was not a major change in the provenance of building blocks when comparing the pre-giant impact Earth, to the inferred composition of the Moon-forming giant impactor. This conclusion is based on the reasoning that the isotopic similarities between the Earth and enstatite chondrites, for lithophile elements such as O and Cr, indicate genetic similarity to enstatite chondrites prior to the giant impact. Yet, the isotopic composition of the siderophile element Mo in Earth’s mantle is also very similar to that of enstatite chondrites. The isotopic composition of Mo present in the mantle was likely strongly affected by additions from the Moon-forming giant impactor, whereas the isotopic compositions of lithophile elements such as O and Cr were not. Thus, the collective enstatite-chondrite-like isotopic compositions of lithophile and siderophile elements suggest that both the Earth and the giant impactor formed in the same region of the protoplanetary disk as enstatite chondrites (e.g., Dauphas et al., 2014).

The genetic heritage of late accreted materials during the final 0.5 wt.% of terrestrial accretion is best monitored via Ru isotopes, combined with the relative abundances of the HSE in the BSE. The Ru isotopic composition of the mantle (Birmingham et al., 2015), as for Mo isotopes, is similar to enstatite chondrites, meaning that the Earth plots near enstatite chondrites at the end of the cosmic Ru–Mo correlation trend. Some aspects of the projected relative abundances of the HSE in the BSE also match certain enstatite chondrites at the end of the cosmic Ru–Mo correlation trend. However, other aspects of the HSE signature of the BSE, such as Ru/Ir, do not match any known chondrite groups. Thus, the late accreted materials must include at least one component with more fractionated HSE than is known to occur in chondrites. The origin of this chemical signature remains poorly constrained, but is suggestive of a not yet sampled primitive meteorite component. Very limited Os isotopic data for Mars suggest a similar late accretionary component was added to its mantle.

Finally, the Earth and Moon were bombarded by an additional flux of planetesimals hundreds of millions of years after primary accretion. The accretionary additions associated with this period could have totaled as much as 0.05 wt.% of the mass of the Earth. The materials involved in this final, minor accretionary period also involved the addition of HSE with some fractionated ratios. The chemical and isotopic natures of

Fig. 13. a–d. Plots of a. Ru/Ir, b. Pt/Ir, c. Pd/Ir, and d. Os/Ir ratios versus $^{187}\text{Os}/^{188}\text{Os}$ (proxy for long term Re/Os) for impact melt rocks from Apollo 14, 15, 16 and 17 landing sites, as well as two lunar meteorites. Each datum represents the regression of data for multiple fragments of each rock (e.g., Fig. 12 for sample 14221). The error bars on the horizontal axes reflect the range of $^{187}\text{Os}/^{188}\text{Os}$ among subsamples of each rock. Data are from: Puchtel et al. (2008); Fischer-Gödde and Becker (2012); Sharp et al. (2014) and Liu et al. (2015). Open symbols are data for bulk chondrites as shown in Fig. 5. Figure is modified from Liu et al. (2015).
these materials are best monitored through the analysis of lunar impact melt rocks that were created by the late-stage basin-forming events. The fingerprinting of these impactors are complex and encompass a range of HSE compositions. Some components evident in this bombardment cohort appear to be similar in HSE characteristics to carbonaceous and ordinary chondrites. An additional component was characterized by substantially higher Re/Os, Ru/Ir and Pd/Ir, compared to any known chondrites, including enstatite chondrites. The dominant signature of at least some materials involved in the LHB, therefore, appear distinct from the prior late-stage building blocks. The Ru and Mo isotopic compositions of lunar impact melt rocks have not yet been determined, so it remains unknown whether or not the LHB can be genetically linked to a type of primitive meteorite.

Although the siderophile element data for the Earth suggest that there was no major change in the provenance of its building blocks through to the end of late accretion (but before LHB), it remains unknown whether or not the building blocks consisted of a genetically homogeneous flux, or included diverse materials that ultimately mixed to form what now appears to be a uniform fingerprint for the BSE. The possibility of isotopic variability of Ru and Mo among potential late stage building blocks, combined with the apparent sluggishness of early mantle mixing of primordial 182W isotopic heterogeneities, suggests that isotopic evidence for diverse late stage impactors might be found in Earth’s early rock record, and possibly in younger rocks. Conversely, given the high level of precision that is now available to search for such isotopic heterogeneities, the future lack of discovery of isotopic anomalies may signal either that the materials involved in the final stages of terrestrial accretion were genetically similar, or that early mixing processes attenuated early Earth heterogeneities before evidence for them could be incorporated in the surviving rock record.

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