Earth and Mars – distinct inner solar system products

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Abstract

Composition of terrestrial planets records planetary accretion, core-mantle and crust-mantle differentiation, and surface processes. Here we compare the compositional models of Earth and Mars to reveal their characteristics and formation processes. Earth and Mars are equally enriched in refractory elements (1.9 × CI), although Earth is more volatile-depleted and less oxidized than Mars. Their chemical compositions were established by nebular fractionation, with negligible contributions from post-accretionary losses of moderately volatile elements. The degree of planetary volatile element depletion might correlate with the abundances of chondrules in the accreted materials, planetary size, and their accretion timescale, which provides insights into composition and origin of Mercury, Venus, the Moon-forming giant impactor, and the proto-Earth. During its formation before and after the nebular disk’s lifetime, the Earth likely accreted more chondrules and less matrix-like materials than Mars and chondritic asteroids, establishing its marked volatile depletion. A giant impact of an oxidized, differentiated Mars-like (i.e., composition and mass) body into a volatile-depleted, reduced proto-Earth produced a Moon-forming debris ring with mostly a proto-Earth’s mantle composition. Chalcophile and some siderophile elements in the silicate Earth added by the Mars-like impactor were extracted into the core by a sulfide melt (~0.5% of the mass of the Earth’s mantle). In contrast, the composition of Mars indicates its rapid accretion of lesser amounts of chondrules under nearly uniform oxidizing conditions. Mars’ rapid cooling and early loss of its dynamo likely led to the absence of plate tectonics and surface water, and the present-day low surface heat flux. These similarities and differences between the Earth and Mars made the former habitable and the other inhospitable to uninhabitable.

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1 Introduction

Earth and Mars share many chemical and physical attributes, but are distinct in size and inventory of volatile elements. Chemical data from surface rocks and meteorites combined with seismological and geodetic observations (e.g., mass, density, moment of inertia (MOI)) provide a multiply constrained compositional model of the terrestrial planets (e.g., Ringwood [1966], Morgan and Anders [1980], Wänke [1981], Longhi et al. [1992]). The physicochemical similarities and differences between Earth and Mars provide useful insights on formation and evolution processes of these bodies, and potentially provide insights into the present-day properties and origin of other rocky bodies (Venus, Mercury, and exoplanets).

By definition, Earth and Mars are located within a habitable zone in which liquid water is available (Cockell et al., 2016; Ehlmann et al., 2016). However, currently life exists only on Earth, and it remains unclear if Mars were inhabited or uninhabited in its history. The occurrence of life only on Earth indicates that there are compositional limits for a habitable planet formation, which can be revealed by an Earth-Mars comparison. Previous comparative studies of chemical compositions of the interiors of Earth and Mars described their differences including abundances of volatile or siderophile elements and redox states (Anders and Owen [1977], Dreibus and Wänke [1987], Wänke and Dreibus [1994]), but the recent advances in compositional modeling of both Earth and Mars will provide further insights into their characters and origins.

Isotopic compositions of solar system materials provide strong constraints on the sources of the terrestrial planets. Mass-independent, nucleosynthetic isotope anomalies in meteorites reveal a heterogeneous distribution of distinct presolar materials and provide the basis for classifying the non-carbonaceous (NC) and carbonaceous meteorite (CC) groups (Trinquier et al., 2007, 2009, Warren [2011], Kruijer et al. [2017a]). The NC meteorites appear to be from the inner solar system, whereas the CC meteorites are considered to be samples from the outer solar system, including the Trojan asteroids and possibly beyond Jupiter’s orbit (Walsh et al., 2011, Kruijer et al., 2017a). These isotopic observations demonstrate a link between Earth and enstatite chondrites, whereas Mars is thought to be most closely related to ordinary chondrites (e.g., Trinquier et al., 2007, 2009, Javoy et al., 2010, Warren, 2011, Dauphas, 2017).

Here we compare the compositional models of Earth (e.g., McDonough and Sun [1995], Palme and O’Neill [2014]) and Mars (e.g., Wänke and Dreibus [1994], Taylor [2013], Yoshizaki and McDonough [2020]) to clarify their characteristics and formation processes. We explore the nature of the building blocks of the terrestrial planets, based on their compositional similarities and differences with chondritic meteorites. These comparisons provide a basis for insights into the conditions for habitable planet formation and evolution.
2 Comparisons

2.1 Bulk planet

The geochemical classification of elements consists of four element groups: lithophile (rock-loving), siderophile (metal-loving), chalcophile (sulfide-loving), and atmophile elements. Lithophile elements (e.g., Si, Mg, Ca, Al, Ti, Na, K, rare earth elements) are preferentially incorporated into oxide phases, and thus concentrated in a silicate shell in a differentiated planetary body. Therefore, their abundances in the mantle can be converted to the bulk composition if we also know a mass fraction of a metallic core in a planet. In addition, elements are also classified based on their volatilities in a gas of solar composition at 10 Pa (Lodders, 2003).

Historically, there have been three families of compositional models for the bulk silicate Earth, in terms of the proportions of the refractory elements relative to Mg and Si, which, together with Fe and O, make up ~90% of rocky planets. These models have low (Javoy, 1995; Javoy et al., 2010 [Warren, 2008; Caro and Bourdon, 2010], medium (Ringwood, 1975; Jagoutz et al., 1979; Wänke, 1981; Hart and Zindler, 1986; Allegre et al., 1995; McDonough and Sun, 1995; Palme and O’Neill, 2014), and high (Wasserburg et al., 1963; Turcotte et al., 2001; Turcotte and Schubert, 2014) proportion of refractory elements (McDonough, 2016 and references therein). Here we estimate a primitive (i.e., the least melt-depleted) mantle composition based on compositional trends from basalts and mantle rocks and propose a model with medium refractory element abundance (e.g., McDonough and Sun, 1995; Palme and O’Neill, 2014; see also Section A.1). As summarized in Palme and O’Neill (2014), these geochemical models of Earth are basically similar to each other, and differences between the models do not affect comparison of Earth and Mars discussed in the current paper.

A similar geochemical approach has been applied in constraining compositional models of Mars (Wänke and Dreibus, 1994; Taylor, 2013; Yoshizaki and McDonough, 2020). Here we adopt the compositional model by Yoshizaki and McDonough (2020), which is compositionally and mineralogically similar to those of Wänke and Dreibus (1994) and Taylor (2013) (Fig. A.1; see also Bertka and Fei, 1997; Khan et al., 2018; Smrekar et al., 2019 for Martian mantle mineralogy models), but differ in model development. For example, Wänke and Dreibus (1994) assumed that Mn and more refractory elements (including Fe, Mg and Si) are in chondritic relative abundances in bulk Mars, which is denied by Yoshizaki and McDonough (2020). The details of previous Mars models are summarized by Taylor (2013) and Yoshizaki and McDonough (2020).

By establishing the planet’s budget of the 36 refractory elements, recognizing that ratios of refractory elements (e.g., Ca/Al, Th/U) and Fe/Ni are constant (±15% or better) in the chondritic building blocks (e.g., Wasson and Kallemeyn, 1988; Alexander, 2019a,b), and either knowing the relative mass of the planet’s core or the mantle’s Mg# (atomic ratio of Mg/(Mg + Fe)), limit the range of acceptable Mg/Si (i.e., olivine/pyroxene) values for planetary compositional models (Figs. 1 and 2A). The Fe/Mg vs Fe/Si correlation seen in both the NC and CC chondrites is suggestive of
metal-silicate gradients in the solar system (Figs. 1 and 2B); interestingly, Earth and Mars also follow this correlation.

The approach used here finds that the bulk Earth contains $1.85 \times$ CI abundances for the refractory elements (e.g., [McDonough and Sun 1995] [Palme and O’Neill 2014]), it has an olivine/pyroxene proportion equivalent to that of pyrolite (Ringwood 1966), and is depleted in volatile elements (i.e., sub-CI Rb/Sr, K/U, and S; Fig. 2A; [Gast 1960] [Wasserburg et al. 1964]). Mars also contains $\sim 1.9 \times$ CI abundances for the refractory elements and is depleted in volatile elements, but less so than Earth (Fig. 2B; [Yoshizaki and McDonough 2020]). The net atomic oxygen/(metallic Fe) values of Earth and Mars (3.7 and 8.7, respectively; Table 1) provide a measure of its average oxidation state. These compositional models for Earth and Mars provide a time-integrated perspective for materials available for accretion at 1 and 1.5 AU, respectively. This spatial sampling contrasts with the temporal comparison, as the mean timescales for Mars’ formation ($\tau_{\text{accretion}}^{\text{Mars}} \sim 2$ Myr; Dauphas and Pourmand 2011) differs from that for Earth ($\tau_{\text{accretion}}^{\text{Earth}} \gtrsim 30$ Myr; Kleine et al. 2009).

2.2 Bulk silicate planet

2.2.1 Geochemistry and mineralogy

Chondritic ratios of refractory lithophile elements in the bulk silicate Earth and Mars (BSE and BSM, respectively) are further confirmed and constrained by Lu-Hf, Sm-Nd and La-Ce isotopic systematics of these planetary materials (e.g., [Bouvier et al. 2008] [Burkhardt et al. 2016] [Willig et al. 2020]). The moderately volatile, lithophile elements (lithophile MVE; e.g., alkali metals) are depleted in Earth and Mars when compared to chondritic meteorites, both planets showing comparable correlations with their condensation temperatures (Fig. 3). Earth, with K/U of 14000 and Rb/Sr of 0.032 (i.e., MVE/refractory ratios; cf., 68000 and 0.30 in CI chondrites, respectively), shows such a strong depletion trend in these elements (Fig. 3A). Mars, with a K/U of 20000 and Rb/Sr of 0.068, also shows this trend (Fig. 3B), albeit less depleted than the Earth’s.

Siderophile and chalcophile elements are depleted in the BSE and BSM compared to lithophile elements with similar condensation temperatures, due to their incorporation into the metallic cores (Fig. 4). The abundances of siderophile elements in the BSE and BSM do not show a sub-parallel trend with that defined by lithophiles (Fig. 4). The degree of siderophile and chalcophile element depletion reflects a combination of planetary building block compositions and element fractionation processes during planetary accretion and differentiation (McDonough 2014).

The BSE and BSM both have high concentrations and chondritic relative proportions of highly siderophile elements (HSE: Re, Os, Ir, Pt, Ru, Rh, Pd and Au; Fig. 4), which is at odds with any combination of high-pressure and temperature conditions for the partitioning of these elements into a core forming melt (Brandon et al., 2012; Walker et al., 2015; Day et al., 2016; Tait and Day, 2018). For example, core-mantle equilibration model in Martian interior (Righter et al., 2015)
predicts sub-CI Re/Os and super-CI Ir/Os and Ir/Ru values in silicates, which are inconsistent with the chondritic HSE pattern in the BSM (Brandon et al., 2012; Tait and Day, 2018). In addition, Righter et al. (2015) assumes a high S concentration (>10 wt%) in the Martian core, which is much higher than a value predicted by the Martian volatility trend of lithophile elements (≤7 wt%; Yoshizaki and McDonough, 2020). These shared features of the HSE abundances in Earth and Mars are considered as evidence for late accretion of chondritic materials during the final stage (e.g., after ≥98% accretion) of planetary growth in the inner solar system (Kimura et al., 1974; Brandon et al., 2012; Day et al., 2016). Delivery of these HSE was also accompanied by an addition of volatile gases and fluids (H, C, N and O; Albarede, 2009; Alexander et al., 2012; Marty, 2012), although the late-accreted material to Mars might have been volatile-depleted (Wang and Becker, 2017; Righter et al., 2019).

On the other hand, the BSE is characterized superchondritic Ru/Ir and possibly Pd/Ir, which cannot be easily accommodated by the late addition of volatile-rich materials (Becker et al., 2006). Experimental studies showed that these high ratios in the BSE might reflect P, T, and composition-dependent changes in partitioning behaviors of Ru and Pd, and proposed no need for the late addition of these elements (Righter et al., 2008; Wheeler et al., 2011; Laarenz et al., 2016; Righter et al., 2018). Alternatively, these inconsistencies between the BSE and chondrites show an unrepresentative sampling of asteroidal materials in our meteorite collections (Walker et al., 2015).

The average, time-integrated, planet-scale redox condition (i.e., metal-silicate equilibrium) is recorded in the Mg# of its mantle, with the BSE and BSM having an Mg# of ~0.89 and 0.75–0.8, respectively, and its core mass fraction (i.e., Mars ~20% and Earth ~32%; Section 2.3). These attributes document Mars being more oxidized than Earth. In addition, Mars lacks depletion in redox-sensitive, nominally lithophile elements (i.e., V, Cr, and Mn), and its mantle has a lower Hf/W value (~3.5; Dauphas and Pourmand, 2011; Yoshizaki and McDonough, 2020), which contrast with those for Earth (Fig. 4). The more oxidized conditions for the Martian mantle are also indicated by mineralogy, trace element compositions and valence states of Fe and Eu in Martian meteorites. The Martian mantle, however, as recorded in Martian meteorites, shows a heterogeneous redox state (e.g., Herd et al., 2001, 2002; Wadhwa, 2001, 2008; Goodrich et al., 2003; McCanta et al., 2009; Shearer et al., 2011; Righter et al., 2016; Herd, 2019). Oxygen might be one of the light elements in the Earth’s core (Ohtani and Ringwood, 1984), but its limited solubility in iron liquids at high P-T conditions (O’Neill et al., 1998) indicates small effects of the core formation in the BSE’s Mg#, and supports the more reduced state of the BSE as compared to the BSM.

Earth and Mars have similar "upper mantle" mineralogies (Fig. 5). Both contain olivine, ortho- and clino-pyroxenes, and garnet, with the Earth’s mantle being richer in olivine than the Martian mantle (~60% and ~45% in modal proportion, respectively). Differences in composition (Table 1) and interior thermal gradient (Breuer and Spohn, 2003; Katsura et al., 2010) of Earth and Mars results in different depths of the olivine-wadsleyite and wadsleyite-ringwoodite phase transitions. For Earth, these transitions are at 410 km and 520 km depth, respectively, and in Mars, they occur at
1000 km and 1300 km depth, respectively. These phase transitions are sharper in the Earth’s mantle than the Martian mantle, because of the former’s higher Mg#, greater garnet abundance, and hotter temperature (Frost, 2003; Filiberto and Dasgupta, 2011; Putirka, 2016). The Martian mantle might not have a layer of bridgmanite, which is a dominant phase in the Earth’s lower mantle (Fig. 6).

2.2.2 Heat producing elements (HPE)

The rocky planets are powered by both primordial accretion energy (and lesser amounts from core formation) and radiogenic heat by radioactive decays of heat producing elements (HPE: K, Th, U). All three of the HPE are incompatible, lithophile elements that have been excluded from their cores and concentrated into planetary crusts (e.g., Corgne et al., 2007; Blanchard et al., 2017; Wipperfurth et al., 2018), which is beneficial for carrying out remote gamma-ray surveys of planetary surfaces (e.g., Peplowski et al., 2011; Surkov et al., 1987; Prettyman et al., 2015). The Martian crust is thicker (estimated to be 30–60 km thick with an average value of ~50 km; Zuber et al., 2000; McGovern et al., 2002; Wieczorek and Zuber, 2004; Humayun et al., 2013), less enriched in incompatible elements, and contains ~50% of the HPE budget of Mars, whereas Earth has a thinner, more chemically evolved crust, which contains only ~35% of the planet’s HPE budget (Table 1 and Section A.2; Rudnick and Gao, 2014). Mars’ internal heating Rayleigh number of >10^5 (Section A.3) is consistent with a convecting Martian mantle and a dynamically stabilized Tharsis bulge (McKenzie et al., 2002; Kiefer, 2003).

With nearly a factor of 2 greater surface to volume ratio for Mars than Earth, the former cooled much faster (Filiberto and Dasgupta, 2011; Baratoux et al., 2011; 2013; Filiberto, 2017; Breuer and Moore, 2015; Putirka, 2016). Mars’ surface heat flux is 19 ± 1 mW/m^2 (Parro et al., 2017) (cf., Earth’s average is 90 mW/m^2; Jaupart et al., 2015), equivalent to a global heat flux of 2.75 ± 0.15 TW, of which 2.5 TW comes from radioactive decay (Table 1 and Section A.2). Mars’ planetary Urey ratio (Ur: total radioactive heat production relative to total surface heat loss) is 0.92 ± 0.05, which is higher than that estimated for Earth (0.43) and its mantle Ur_Mars (planetary Ur_Mars minus the crustal fraction) is 0.8, more than double of that of the Earth’s mantle (0.33). The high Ur_Mars demonstrates a minor contribution from secular cooling to the total Martian heat flux. Importantly, the modeled Martian surface heat flow varies between 14 and 25 mW/m^2 (Parro et al., 2017). A direct measurement the Martian surface heat flux by NASA’s Interior Exploration using Seismic Investigations, Geodesy and Heat Transport (InSight) mission (Banerdt et al., 2020) will provide important constraints on its heat production and thermal history.

2.3 Core

The mantles of both Earth and Mars are depleted in siderophile and chalcophile elements due to core extraction (Fig. 4). Their core compositions are modeled using the following constraints: the planet’s chondritic Fe/Ni value, temperature, mass, density, MOI, and for Earth, seismic profile (see
McDonough, 2014; Yoshizaki and McDonough, 2020). Also, based on iron meteorite mineralogy (Scott and Krot, 2014) and cosmic abundance of elements (Lodders, 2020), planetary cores might contain sulfides, carbides and phosphides, and possibly other phases (e.g., silicides if formed under highly reduced conditions).

The Earth’s core (32% by mass) has an outer liquid layer and a solid inner core (∼5 wt%) (Fig. 6). Estimates of the size of the Martian core ranges from 18–25 wt% and 1500–1900 km, respectively (Section A.1.3). Data from multiply orbiting satellites provide a precise (±2%) measure of Mars’ Love number solution \(k_2\) and document a possible existence of a partially molten core (Yoder et al., 2003; Genova et al., 2016; Konopliv et al., 2016). This observation indicates significant amounts of light elements in the core that lowers its solidus temperature.

Sulfur is a prime candidate for light elements in a planetary core. The volatile element depletion trends for both Earth and Mars (Fig. 4) constrain the core’s S content of ∼1.8% in Earth and ∼7% in Mars (Yoshizaki and McDonough, 2020; see also Sections A.1.2 and A.1.3). Since the estimated S contents of the Earth’s and Mars’ core are not high enough to explain their respected core density deficits, additional light elements in these cores are needed (e.g., Si, O, H and C). The additional light elements are likely to be different in these cores because metal-silicate partitioning behavior of elements depends on conditions of core formation (e.g., timing, \(P\), \(T\), \(f_{O_2}\)) and compositions of coexisting silicate and metallic phases.

As described above, a planet’s volatility trend sets expectations for the absolute amount of siderophile and chalcophile elements in the planet and defines the proportion of these elements that were partitioned into the core. The degree of depletion below the volatility trend defines an element’s empirically established metal/silicate partition coefficient (McDonough, 2014; Yoshizaki and McDonough, 2020). The Martian core is enriched in siderophile and chalcophile MVE (e.g., P, Ge, S) as compared to the Earth’s, and there appears to be little to no Ga in the cores of terrestrial planets, which contrasts with that seen in iron meteorite compositions (Figs. 7 and 8). Compared to the Earth’s core, the Martian core is smaller, it might contain larger amounts of H and O, and it has a lower Fe/Ni value, with its mantle being enriched in Fe (Table 1). Overall, these factors contribute to Mars’ lower uncompressed density as compared to the Earth’s (3750 vs 4060 kg/m\(^3\); Table 1). There is no unique model for core compositions of Earth and Mars that satisfies the geodetic and geochemical constraints (McDonough, 2014; Yoshizaki and McDonough, 2020), and further experimental and theoretical efforts are needed to constrain the light element budget in the planetary cores.
3 Discussion

3.1 Origin of volatile depletion in terrestrial planets

From Mercury to Mars and beyond to Vesta, the second biggest asteroid, their surface K/Th values are significantly lower relative to chondrites (Surkov et al., 1987; Taylor et al., 2006; Peplowski et al., 2011; Prettyman et al., 2015). The origins of this depletion and that of other MVE remain elusive, with possible explanations including post-nebular volatile loss due to internal or impact-induced heating (e.g., O’Neill and Palme, 2008; Norris and Wood, 2017) and incomplete condensation of nebular gas (e.g., Palme and O’Neill, 2014; Siebert et al., 2018). Since planetary K/Th ratios and relative amounts of more refractory olivine (Mg$_2$SiO$_4$) and less refractory pyroxene (MgSiO$_3$) do not correlate with their heliocentric distances in both the solar and extra-solar systems (e.g., Fig. 2A; van Boekel et al., 2004; Kessler-Silacci et al., 2006; Bouwman et al., 2008, 2010; Sargent et al., 2008), these volatile depletions do not solely reflect an outward temperature decrease in the protoplanetary disk.

The absence of heavy isotope enrichment in rocks from Earth and Mars for multiple isotope systems (e.g., K, Zn, Rb, Fe, Cd; Humayun and Clayton, 1995; Nebel et al., 2011; Paniello et al., 2012a; Sossi et al., 2016, 2018; Pringle and Mynier, 2017; Wombacher et al., 2008) indicates negligible post-accretionary evaporative loss of the MVE. In contrast, isotopically heavier siderophile or chalcophile compositions (e.g., Fe, Ga, Sn) of terrestrial and lunar mantles might reflect (1) core-mantle differentiation, (2) evaporation of the giant impactor’s core and addition of the metal/sulfur-loving elements to the terrestrial mantle during the giant impact (see Section 3.3), or (3) planetary surface processes (e.g., Poitrasson et al., 2004; Kato and Mynier, 2017; Creech and Moynier, 2019). The $^{53}$Mn-$^{53}$Cr and $^{87}$Rb-$^{87}$Sr isotope systematics of meteorites and terrestrial samples are consistent with a volatility-dependent, gas-solid fractionation during first few Myr of the solar system (e.g., Shukolyukov and Lugmair, 2006; Trinquier et al., 2008; Hans et al., 2013; Mynier et al., 2012), which might be prior to the dissipation of the nebular gas ($\sim t_0 + 5$ Myr; Wang et al., 2017).

There is negligible evidence for post-accretionary MVE losses in the chemical compositions of these planets. The bulk Earth and Mars, together with chondrites, show negligible evidence of evaporative losses in ratios of Na, Mn and Ti (Fig. 9 and Section A.1.1; O’Neill and Palme, 2008; Siebert et al., 2018), which have distinct relative volatilities during condensation and evaporation. Values of Mn/Na and Na/Ti in the bulk Earth, Mars and chondrites are consistent with an incomplete nebular condensation, in which earlier condensates are removed from a nebular gas before a completion of more volatile species (e.g., Larimer, 1967; Larimer and Anders, 1967) (cf. O’Neill and Palme, 2008). The plot of Mn/Na versus Na/Ti indicates that a formation of precursors of Earth and Mars at higher temperatures than their isotopically linked counterparts (enstatite and ordinary chondrites, respectively), which is also suggested by the former’s higher Mg/Si (i.e., olivine/pyroxene), Al/Si, and Al/Mg ratios (Kerridge, 1979; Larimer, 1979; Dauphas et al., 2015; Morbidelli et al., 2020).

Collectively, the compositions of rocky planets, as compared to their chondritic relatives, likely
reflects volatility-dependent chemical fractionation in the protoplanetary disk, rather than the post-accretionary losses of MVE. In contrast, small differentiated asteroids (e.g., parent bodies of Eucrite and Angrite) show clear evidence for preferential loss of MVE (Fig. 9; O’Neill and Palme, 2008) and heavy isotope enrichment (e.g., Paniello et al., 2012b; Pringle et al., 2017; Tian et al., 2019), which are the hallmarks of evaporative losses during or after accretion. Volatile elements might have escaped from these small bodies during and/or after their accretion, due to their weak gravity field.

3.2 The building blocks of the terrestrial planets

The planetary building blocks appeared to be made up of high-temperature materials, dominantly chondrules, which are silicate droplets formed by transient heating events within first few Myr of the solar system evolution (Connelly et al., 2012; Bollard et al., 2017), and are an essential component of chondrites (e.g., Scott and Krot, 2014; Russell et al., 2018). MVE composition of chondrules provides a record of incomplete nebular condensation rather than evaporation processes (Humayun and Clayton, 1995; Alexander et al., 2000, 2008; Galy et al., 2000; Pringle et al., 2017). The refractory element enrichment and MVE depletion in the BSE and BSM are comparable to those observed for chondrules from carbonaceous chondrites (Fig. 10; Hewins and Herzberg, 1996; Mahan et al., 2018). Significantly, mass-dependent Ca (Huang et al., 2010; Magna et al., 2015; Amsellem et al., 2017; Simon et al., 2017; Birmingham et al., 2018) and Mg (Bizzarro et al., 2004; Young and Galy, 2004; Wiechert and Halliday, 2007; Bouvier et al., 2013; Olsen et al., 2016; Hin et al., 2017) isotopic compositions of chondrules, BSE and BSM also suggest that they inherited fractionated isotopic signatures from similar precursor materials, which have experienced high-temperature gas-solid fractionation processes.

Recent theoretical models of planetary growth favor formation of terrestrial planets via accretion of chondrule-sized pebbles (e.g., Johansen et al., 2015a,b; Levison et al., 2015). These models predict a rapid accretion of Mars-sized bodies under the presence of a nebular disk. Thus, Mars, with a $\tau_{\text{accretion}}^{\text{Mars}} \sim 2$ Myr (Dauphas and Pourmand, 2011; Kruijer et al., 2017b; Bouvier et al., 2018) (cf. Marchi et al., 2020), formed within the nebular disk lifetime ($\sim 5$ Myr; Wang et al., 2017), whereas Earth is suggested to have $\tau_{\text{accretion}}^{\text{Earth}} \geq 30$ Myr (e.g., Kleine et al., 2009), which documents its accretion stretched beyond the lifetime of the nebular disk.

Differences in planetary MVE abundances (Fig. A.3) likely reflect aspects of their accretion histories and sizes. Chondrites, the least MVE-depleted bodies’ materials, contain fine-grained, volatile-rich matrix that accounts for the largest fraction of the inventory of volatiles (e.g., Alexander, 2005, 2019a,b; Bland et al., 2005; Zanda et al., 2018). Although a relationship between chondrules and matrix remains poorly understood, their coexistence in the protoplanetary disk before planetesimal accretion is accepted (e.g., Hezel et al., 2018a; Zanda et al., 2018). Some carbonaceous chondritic asteroids accreted more chondrules and less matrix, resulting in more MVE-depleted compositions (Figs. 11B and A.4). According to the pebble accretion model (Johansen et al., 2015a,b), the rapid accretion of pebbles (e.g., Bollard et al., 2017) under the presence of a nebular disk forms terrestrial planets with $\tau_{\text{accretion}} \leq 2$ Myr (Dauphas and Pourmand, 2011; Kruijer et al., 2017b; Bouvier et al., 2018) (cf. Marchi et al., 2020), whereas Earth is suggested to have $\tau_{\text{accretion}} \geq 30$ Myr (e.g., Kleine et al., 2009), which documents its accretion stretched beyond the lifetime of the nebular disk.
(Levison et al., 2015), planetesimal growth starts off by preferentially accreting the smallest particles, and as the body grows, it prefers to accrete larger and larger size particles. This mechanism would lead to a growing planetesimal having a larger chondrule/matrix ratio, and becoming more MVE-depleted as its mass increases. Thus, small chondritic asteroids likely co-accreted chondrules and matrix, whereas Mars, with its intermediate volatile depletion, size and accretion timescale, likely accreted a greater fraction of chondrules to matrix (Fig. 11). Finally, the Earth’s prolonged accretion history was dominated by chondrule accretion, resulting in its significant MVE-depleted composition. The MVE depletion scales with size of the chondritic parent body and planet (Fig. 11), implying an accretion driven process from an undepleted nebula (i.e., CI (solar) composition). Exceptions are small differentiated bodies (e.g., the Moon and Vesta) that experienced post-nebular volatile loss due to internal or impact-induced heating (Section 3.1). Thus, the chondrule-driven planetary growth plays a critical role in establishing the planetary MVE-depleted characteristics.

In contrast, enstatite and ordinary chondrites, whose isotopic composition and redox state are most related to Earth and Mars, respectively (e.g., Trinquier et al., 2007, 2009; Warren, 2011; Dauphas, 2017, Fig. 2B), do not show a clear refractory enrichment nor MVE depletion (Figs. 2A, 11 and A.4). In addition, chondrules from these NC chondrites are less depleted in MVE than the BSE and BSM (Fig. A.5). Thus, the chondrule-driven MVE-depletion scenario discussed above cannot be applied to the NC chondrites we have in our collections. These observations indicate that the NC chondrites represent refractory-poor, volatile-rich counterparts of the inner rocky planets. Morbidelli et al. (2020) showed that the low Mg/Si and Al/Si solids, which are comparable to those of NC chondrites (Fig. 2A), condense after removal of early-formed, high-temperature condensates from the system.

Differences in the timing of planetary accretion might also be important in establishing their relative abundances of MVE. Chronology of meteorites combined with thermal modeling of asteroids (Sugiura and Fujiya, 2014; Kruijer et al., 2017a; Zhu et al., 2020) indicates that the undifferentiated NC planetesimals accreted 1–2 Myr after formation of differentiated NC bodies (i.e., iron meteorite parent bodies and terrestrial planets). The variation in the absolute ages of chondrules (Connelly et al., 2012; Bollard et al., 2017), occurrence of relict grains (Jones, 1994; Weisberg et al., 2011; Tenner et al., 2018) and igneous rims (Krot and Wasson, 1995) in chondrules, and the presence of compound chondrules (Wasson et al., 1995) indicate that chondrule formation was a repeated event. Thus, the longer chondrules remained in the accretionary disk, the more opportunities it has being recycled by later events. Ordinary chondrite chondrules records an admixing of MVE-rich CC-like materials to MVE-poor chondrule precursors into the NC reservoir (Mahan et al., 2018; Schiller et al., 2018; Bollard et al., 2019). Additions of MVE-rich materials and repeated chondrule recycling produce younger chondrules with higher MVE abundance (Mahan et al., 2018). Thus, terrestrial planets, which are dominated by earlier materials that experienced fewer recycle events and MVE addition, are more depleted in MVE as compared to the NC chondrites, which accreted the younger MVE-enriched chondrules.

In contrast, chondrule formation scenarios predict less frequent chondrule formation/recycle
events in the outer solar system (e.g., Morris et al., 2012; Johnson et al., 2015; Sanders and Scott, 2018; Pilipp et al., 1998). This prediction is consistent with chemical and isotopic signatures of CC chondrules (e.g., Hewins and Zanda, 2012; Tenner et al., 2018; Mahan et al., 2018). Consequently, the CC chondrules could have preserved their MVE depletion until the accretion of CC chondritic asteroids, and thus are chemically comparable to the early NC materials which formed the inner rocky planets.

This chondrule-rich accretion model for the terrestrial planets reveals the limited ability to reach greater levels of enrichment in refractory elements (Figs. 2A and 10B), which is a concern for planetary compositional models promoting high refractory element abundance (Section 2.1). To reach higher levels of refractory element enrichment beyond that seen in CV chondrites and their chondrules, larger amounts of refractory inclusions are needed, which is inconsistent with chondritic REE ratios of the BSE (Stracke et al., 2012; Dauphas and Pourmand, 2015).

3.3 Formation models of the terrestrial planets

3.3.1 Previous models of the Earth’s accretion and Moon formation

Depletion of moderately to highly siderophile and chalcophile elements in the BSE is consistent with Earth’s initial accretion from highly reduced, volatile-depleted materials, that was later oxidized by volatile-rich additions (e.g., Wänke, 1981; Wänke et al., 1984; Wänke and Dreibus, 1988; O’Neill, 1991a, 1991b; Rubie et al., 2011, 2015). Support for this temporal evolution in volatile accretion is found in multiple radiogenic isotope systems (e.g., Rb-Sr, U-Pb, Ag-Pd, I-Pu-Xe; Halliday and Porcelli, 2001; Albarede, 2009; Schönbachler et al., 2010; Mukhopadhyay, 2012; Ballhaus et al., 2013; Maltese and Mezger, 2020), metal-silicate partitioning behaviors of elements such as W, Mo, S and C (e.g., Wade et al., 2012; Li et al., 2016; Suer et al., 2017; Tsuno et al., 2018; Ballhaus et al., 2017), and N-body simulations of planetary formation (e.g., Raymond et al., 2006; Morbidelli et al., 2012). However, the degree of volatile element depletion in the pre-impact proto-Earth remains unconstrained.

The two-component accretion models for the growth of Earth envisage mixing of highly reduced, volatile-depleted materials with oxidized, volatile-rich, “CI-chondritic” materials in some proportion, not well defined, but generally conceived their mass ratios to be in the range of 60:40 to 90:10 (Wänke et al., 1984; Wänke and Dreibus, 1988; O’Neill, 1991b). Some mass estimates of the late oxidized addition are comparable to a Mars-sized, Moon-forming impactor (e.g., Canup and Asphaug, 2001; Canup, 2004, 2008). The impactor’s composition has often been assumed to be CI-chondritic (e.g., O’Neill, 1991a,b; Maltese and Mezger, 2020).

However, the CI-chondritic impactor model has been multiply challenged. The common MVE depletion in rocky differentiated bodies in the solar system (e.g., sub-solar K/Th ratios; Section 3.1) might indicate a similar MVE depletion in the Mars-sized impactor. In addition, the CI-like impactor model requires a significant volatile depletion in the proto-Earth, perhaps at levels seen in
angrites and calcium-aluminum-rich inclusions (CAIs), which show heavy Mg- and Si-isotope enrichment and significant depletion of moderately volatile elements (e.g., Grossman et al., 2000, 2008; Pringle et al., 2014) due to significant evaporative losses of the major and more volatile elements by impact-induced or transient nebular heating events (e.g., Stolper and Paque, 1986; Richter et al., 2002; Pringle et al., 2014; Yoshiizaki et al., 2019; Young et al., 2019). However, such isotopic signatures are not recognized for Earth (Section 3.1). Furthermore, models predicting compositional zoning in the protoplanetary disk have a region of CI-like material accreting at \( \geq 15 \) AU (Desch et al., 2018), where it is predicted that the disk mass is too low to form a Mars-sized body.

Compositional similarities of Earth and Moon in multiple isotope systems set strict constraints on the nature of both the impactor and proto-Earth, that is, they are derived from a similar NC isotopic reservoir that appears to be restricted to inner solar system sources (e.g., Wiechert et al., 2001; Trinquier et al., 2009; Warren, 2011; Zhang et al., 2012; Greenwood et al., 2018; Dauphas, 2017; Kruijer et al., 2017a). This observation is consistent with dynamical simulations which predict low probabilities of a Moon-forming impactor originating from \( > 10 \) AU (e.g., Jackson et al., 2018). Thus, these constraints exclude the isotopically distinct CI chondrite as an impactor candidate. In contrast, recent Mo and O isotopic data from a broad range of NC and CC materials challenge this exclusion and find support for a CC-like impactor and/or vigorous mixing of proto-Earth and impactor, requesting further constraints to reveal the origin of the Earth-Moon system (Young et al., 2016; Budde et al., 2019; Cano et al., 2020). Furthermore, the recently-proposed synestia model (Lock et al., 2018) has the Moon forming in a vapor cloud surrounding Earth. This vapor cloud was produced by a large impact, resulting in a well-mixed, chemically equilibrated proto-earth and the vapor cloud.

### 3.3.2 Mars-like Moon-forming giant impactor model

Here we propose a model for the origin of the Moon. Our model is a modification of the original Wänke et al. (1984)’s model and later modified by O’Neill (1991a,b). In our version of this model, we envision the Earth’s formational history in 4 steps and use a differentiated Mars-like composition (Yoshizaki and McDonough, 2020), instead of CI composition, for the giant impactor. Figs. 12 and 13 show the details of our model: Fig. 12 specifically highlights the differences between our model and those previously presented. Our model starts with

1. the accretion of the proto-Earth (reduced and volatile-depleted) accompanied by continuous core-mantle differentiation (\( \sim 90\% \) of Earth’s mass),

2. followed by a late-stage Moon-forming giant impact event (30–100 Myr after \( t_0 \); Kleine et al., 2009) that adds the final \( \sim 10\% \) mass (oxidized and volatile-enriched) to Earth and forms a protolunar accretion disk,

3. subsequently, the mantle loses a Fe-Ni (\( \pm O \)) sulfide liquid (sulfide matte; O’Neill, 1991b) to the core (\( \sim 0.5\% \) BSE mass),
4. and finally, the BSE receives the addition of (∼0.5% of the BSE mass) a late accretion component that brings the highly siderophile and chalcophile elements in chondritic proportions and highly volatile gases and fluids (see Section A.5 for details of the modeling).

In this scenario, the bulk proto-Earth already contains ≥80% of the present-day Earth’s budgets of most of the MVE (e.g., K/Th ∼ 3200; Rb/Sr ∼ 0.026; Table A.2), since a Mars-like impactor contributes only a limited amount of additional MVE (Fig. 13). The addition of MVE by the impactor leads to sulfur saturation in the magma ocean, generating an Fe-Ni (± O) sulfide liquid (post-impact sulfide matte; O’Neill, 1991b; Rubie et al., 2016). The sulfide matte precipitates through a crystallizing mantle into the core due to its high immiscibility, low wetting angle, and high density (Gaetani and Grove, 1999; Rose and Brenan, 2001). Assuming that the sulfide matte, with a present-day mantle sulfide-like composition (Lorand and Conquéré, 1983), extracted all S from the post-impact Earth’s mantle; its mass is estimated to be ∼0.5 wt% of the present-day Earth’s silicate mantle (Section A.5).

The siderophile and chalcophile element abundances of the BSE are reproduced when 10–15% of a planetary mass is added by the impactor and most of the impactor’s core equilibrates with the proto-Earth’s mantle (Fig. 13 and Section A.5). In this scenario, the bulk proto-Earth contains ≥80% of the present-day Earth’s budget of most of the MVE (e.g., K/Th ∼ 3200; Rb/Sr ∼ 0.026; Table A.2), with the Mars-like impactor contributing a limited amount of MVE (Fig. 13). We envision the proto-Earth as having a MVE abundance comparable to that of chondrules (Figs. 10 and 11B). Thus, the proto-Earth’s composition is consistent with the chondrule-rich accretion scenario (Section 3.2) and requires no need for a post-accretionary loss of MVE from the proto-Earth before the Moon-forming event (Section 3.1).

The mass fraction of the impactor contributing to the lunar composition provides a critical constraint on the lunar formation models. The lunar mantle is depleted in nominally lithophile V, Cr and Mn (Dreibus and Wänke, 1979). Such lunar mantle depletion can be achieved by high-\(T\) or S-rich conditions during lunar core formation, but these conditions seem unlikely (Steenstra et al., 2016). If a Mars-sized impactor with no V, Cr or Mn-depletion in its mantle contributed >70% of Moon, as predicted by the canonical giant impact models (Canup and Asphaug, 2001; Canup, 2004, 2008), ≥40% of evaporative losses of Mn and Cr are needed to produce their depletion in the lunar mantle (Fig. 14). Such significant evaporation of these elements is inconsistent with their least volatile nature among MVE (Gellissen et al., 2019; Sossi et al., 2019). Thus, the proto-Earth’s V, Cr and Mn-depleted mantle should be a primary source of the Moon-forming materials, as supported by their isotopic similarities (e.g., Wiechert et al., 2001; Warren, 2011; Zhang et al., 2012; Greenwood et al., 2018). A recent geochemical model of Earth’s Moon formation and differentiation prefers a present-day BSE-like MVE composition for the bulk proto-Moon (i.e., lunar source materials before gas-melt fractionation; Righter, 2019), which is consistent with the proto-Earth’s MVE abundance and its large contribution to the lunar source materials (Fig. 14 and Table A.2). The proto-Earth origin of the Moon is also consistent with recent particle hydrodynamic collision
simulations (Hosono et al. 2019) which showed that a terrestrial magma ocean might be a source of lunar building blocks.

### 3.3.3 Mars’ accretion and core formation

Our compositional estimate of Mars requires a much simpler formation history as compared to Earth’s multi-stage formation scenario. The lack of depletion in nominally lithophile elements in the BSM (V, Cr, and Mn; Fig. 4) is consistent with its accretion under uniformly oxidized and low-\(T\) conditions (e.g., Wood et al. 2009). Mars might have experienced a large impact(s) that produced its hemispheric crustal dichotomy and possibly Martian moons (e.g., Marinova et al. 2008; Canup and Salmon 2018), but estimates of a putative small impactor (\(\sim 0.1\%\) of the Mars’ mass; Canup and Salmon 2018) would result in negligible changes in its bulk composition. The 0.5–1\% mass addition of chondritic late accretion material provides highly siderophile and chalcophile elements to the Martian mantle (Tait and Day 2018; Yoshizaki and McDonough 2020).

Righter and Chabot (2011) and Yang et al. (2015) estimated Martian core formation happening at 14 ± 3 GPa and 2100 ± 200 K based on its previous compositional models. Following the approaches of Righter and Chabot (2011) and Yang et al. (2015), the Martian siderophile element distribution of Yoshizaki and McDonough (2020) can be modeled by single stage \(P\)-\(T\) and redox conditions (Fig. A.8), whereas there is no unique solution for Martian core formation (Fig. A.9).

The simple formation history of Mars, combined with its rapid and early accretion (Dauphas and Pourmand 2011; Kruijer et al. 2017b; Bouvier et al. 2018; Marchi et al. 2020), is consistent with its status as a planetary embryo. Given compositional and redox state gradients in the protoplanetary disk, Mars might record the chemical characteristics of nebular materials in the Mars’ orbit in the first few years of the solar system, whereas Earth might have incorporated oxidized materials from greater heliocentric distance in its later accretion stage (Section 3.3.2; Rubie et al. 2015). The Mars’ status as a planetary embryo suggest that a Mars-sized body commonly has a Mars-like composition, supporting the Mars-like Moon-forming impactor scenario (Section 3.3.2).

### 3.4 Conditions for a habitable planet formation

The early \(\tau_{\text{accretion}}\) age implies that Mars underwent global-scale silicate melting and rapid core formation due to heating from short-lived \(^{26}\text{Al}\) and \(^{60}\text{Fe}\) (Fig. A.12; Dauphas and Pourmand 2011; McDonough et al. 2020). The peak radiogenic heating occurs at about 1 to 5 Myr after \(t_0\), well within the time scale for Mars accretion. With \(\tau_{\text{accretion}} = 2\) Myr, the radiogenic energy supplied by \(^{26}\text{Al}\) is comparable to Mars’ gravitational binding energy (\(\sim 7 \times 10^{29}\) J and \(\sim 10^{30}\) J, respectively). During the first 10 Myr, radiogenic heating (\(^{26}\text{Al}, ^{60}\text{Fe}\) and \(^{40}\text{K}\), in order of significance) is comparable to the planet’s primordial energy and is a major control on its thermal evolution.
Mars has the attributes needed for a rocky planet to be biologically available in its early history (e.g., Ehlmann et al., 2016), and it has a higher bulk heat production than Earth’s (3.9 vs 3.3 pW/kg; Table 1). Nonetheless, it has rapidly lost much of its primordial energy (i.e., accretion and core differentiation) due to its larger surface to volume ratio (a factor of 2) and smaller core size (i.e., reduced bottom heating), and it is in waning stages of limited fuel resources (Parro et al., 2017; Yoshizaki and McDonough, 2020). Basal heating of the Martian mantle by its core enhances its thermal evolution, while the transfer of hydrogen from the adjacent ringwoodite to the core (Shibazaki et al., 2009; Yoshizaki and McDonough, 2020) reduces the lifetime of the dynamo (O’Rourke and Shim, 2019). Collectively, these processes likely contributed to dynamo termination at ~4 Ga and loss of the protective magnetosphere (Acuña et al., 1999; Arkani-Hamed, 2004; Lillis et al., 2008). This magnetosphere shields the planet from atmospheric losses, enhances its surface UV radiation, and leads to dramatic climate changes (Ehlmann et al., 2016).

Volatile elements may play a critical role in establishing the amount of light elements in and solidus of the core. The amount of water and other volatile species in the planet’s interior and surface may potentially create the appropriate conditions for the initiation of plate tectonics (e.g., Albarede, 2009; Ehlmann et al., 2016). Likewise, the heat-producing elements and a reduced core solidus keep the metallic core convecting and lead to the creation of a magnetic field, which shields a planet’s surface from cosmic rays. Together, heat-producing and volatile elements regulate a planet’s cooling history, drive crustal differentiation, and make it habitable (Ehlmann et al., 2016). The simple formation history of Mars (Section 3.3.3; Rubie et al., 2015) emphasizes the uniqueness of Earth, the sole habitable planet in our solar system. In turn, the Earth-Mars comparison indicates that high-temperature nebular chemical processes and timescales of planetary accretion are essential in making habitable planets.

Depletion in volatile elements appears to be a common feature of the terrestrial planets (Fig. 3; Surkov et al., 1987; Peplowski et al., 2011), and may likely be so for rocky exoplanets (Harrison et al., 2018, 2021). The relationship between the planetary volatile depletion, size, accretion timescale and abundance of chondrules (Fig. 11) predict accretion timescales of Venus and Mercury of 30–100 Myr and 2–10 Myr, respectively, based on their Earth- and Mars-like K/Th ratios, respectively (Surkov et al., 1987; Peplowski et al., 2011). The predicted ages of Mercury and Venus provide a foundation for future investigation of their thermal history and habitability.

Venus and Earth are quite similar in their physical and chemical properties (size, bulk K/Th ratio, Mg# in basalt and possibly MOI; e.g., Surkov et al., 1987; Dumoulin et al., 2017), but their present-day surface conditions are distinct. Venus does not show a clear evidence for a giant impact, and its size and K/Th ratio (~3000; Surkov et al., 1987) are comparable to those of the proto-Earth (K/Th ~ 3200; Table A2). Therefore, the present-day Venus might be comparable to the pre-impact proto-Earth. Further observational, cosmochemical and theoretical investigations of Venus may provide useful insights into the pre- and post-formational history of Earth, the only habitable planet in the today’s solar system.
4 Conclusions

The refractory element enhancement and volatile depletion of Earth and Mars were established by a nebular chemical fractionation. Post-accretionary losses of moderately volatile elements are negligible. The degree of volatile element depletion correlates with the abundance of accreted chondrules, planetary size, and their accretion timescale. The present-day bulk silicate Earth composition is consistent with a Moon-forming impactor having a Mars-like size and composition. Planetary chemistry, which is related to many factors including the building block composition, the timing, duration and sequence of accretion and its differentiation history, play an essential role in making a planet habitable.

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Author contributions

TY and WFM proposed and conceived various portions of this study and together calculated the compositional models of planets. The manuscript was jointly written by TY and WFM and they read and approved the final manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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Data and materials availability

Materials used in this study are available within the paper or supplementary materials.
Table 1: Physical and chemical properties of Earth and Mars. Modeled values are in normal and reference values in italic fonts.

| Observation | Unit     | Crust     | Mantle    | Core       | Bulk planet  | Reference value |
|-------------|----------|-----------|-----------|------------|--------------|-----------------|
| **Earth**   |          |           |           |            |              |                 |
| Mass        | kg       | 3.12 × 10^{22} | 4.00 × 10^{24} | 1.94 × 10^{24} | 5.972 × 10^{24} | 5.97218(60) × 10^{24} |
| Mean density | kg/m^3   | 2800      | 4400      | 11870      | 5514         | 5514(2)         |
| Moment of inertia | – | 1% | 88% | 11% | 0.3308 | 0.330690(9) |
| Heat production (K, Th, U) | TW | 7.3 | 12.6 | 0 | 19.9 | 46(3) |
| **Mars**    |          |           |           |            |              |                 |
| Mass        | kg       | 2.56 × 10^{22} | 5.01 × 10^{23} | 1.17 × 10^{23} | 6.419 × 10^{23} | 6.417(3) × 10^{23} |
| Mean density | kg/m^3  | 3010      | 3640      | 6910       | 3936         | 3935(1)         |
| Moment of inertia | – | 7% | 89% | 4% | 0.3638 | 0.3639(1) |
| Heat production (K, Th, U) | TW | 1.3 | 1.3 | 0 | 2.5 | 2.7(2) |

| Observation | Unit     | Crust  | Mantle  | Core     | Bulk planete | Chondrite   |
|-------------|----------|--------|---------|----------|--------------|-------------|
| SiO_2       | wt%      | Earth  | Mars    | Earth    | Mars         |            |
|             |          | 44.9   | 45.5    | 79.5     | 36.3         | 30.0        |
| TiO_2       |          | 0.20   | 0.17    | 7.4      | 32.0         | 31.0        |
| Al_2O_3     |          | 4.44   | 3.59    | 5.2      | 15.4         | 11.3        |
| MnO         |          | 0.14   | 0.37    | 6.6      | 16.1         | 17.4        |
| FeO         |          | 8.06   | 14.7    | 0.9      | 1.82         | 1.4         |
| MgO         |          | 37.8   | 31.0    | 7.4      | 1.87         | 1.27        |
| CaO         |          | 3.54   | 2.88    | 3.3      | 1.71         | 1.38        |
| Na_2O       |          | 0.36   | 0.59    | 0.9      | 0.59         | 0.2         |
| K_2O        |          | 0.034  | 0.043   | 0.17     | 0.07         | 0.2         |
| P_2O_5      |          | 0.021  | 0.17    | 0.046    | 0.07         | 0.2         |
| NiO         |          | 0.25   | 0.046   | Total    | 99.7         | 100.2       |
| Cr_2O_3     |          | 0.15   | 0.88    | Mg/Si    | 0.96         | 0.63        |
| K (ppm)     |          | 280    | 360     | Al/Si    | 0.10         | 0.05        |
| Th (ppb)    |          | 76     | 68      | Fe/Si    | 2.0          | 1.7         |
| U (ppb)     |          | 20     | 18      | Total    | 100.2        | 99.9        |

1Chambat et al. (2010). 2Wipperfurth et al. (2020). 3Dziewonski and Anderson (1981). 4McDonough (2014). 5Jaupart et al. (2015). 6Rudnick and Gao (2014). 7Global surface heat loss. 8Konopliv et al. (2016). 9Rivoldini et al. (2011). 10Taylor and McLennan (2009). 11Yoshizaki and McDonough (2020). 12Parro et al. (2017). 13This study. 14Alexander (2019a,b) (Volatile-free, normalized to total = 100 wt%).
Fig. 1: (A) Ternary plot and (B) scatter ratio plot of major elements Si, Fe and Mg in bulk Earth (McDonough, 2014), bulk Mars (Yoshizaki and McDonough, 2020) and chondritic meteorites (Alexander, 2019a,b). The regression line for chondrites (except for sulfide-rich EH) is also shown in the lower panel. TL—Tagish Lake.
Fig. 2: (A) Weight ratios of Al/Si vs Mg/Si values for planets and chondrites. Earth and Mars show higher RLE abundances (i.e., Al) and Mg/Si value (i.e., olivine/pyroxene ratio) compared to enstatite and ordinary chondrites, which are respectively their isotopically identified relatives. (B) The Urey-Craig diagram (after Urey and Craig, 1953) illustrates relative redox condition for Earth, Mars, and chondrites. Chondrite classification (non-carbonaceous (NC; red) vs carbonaceous (CC; blue) groups) follows Warren (2011) and Kruijer et al. (2017a). TL–Tagish Lake. Data sources are as in Fig. 1.
Fig. 3: Lithophile element abundances in the bulk silicate Earth and Mars normalized to CI abundances and refractory lithophile element Al. The scale of right ordinate shows the CI-normalized abundances. The values are plotted against 50% condensation temperature (K) of elements at 10 Pa (open; Wood et al., 2019) and (filled; Lodders, 2003). CI abundance is from Lodders (2020). Other data sources are as in Fig. [1].
Fig. 4: Siderophile (red square) and chalcophile (green triangle) element abundances in the bulk silicate Earth and Mars plotted against 50% condensation temperature (K) of elements at 10 Pa (Wood et al., 2019). The values for y axes follow the same convention as Fig. 3. Also shown are time-integrated planetary volatility trends at 1 AU (Earth) and 1.5 AU (Mars) defined by lithophile elements (Fig. 3). Data sources are as in Fig. 1.
Fig. 5: Mineralogy and physical properties of planetary mantles. (A, C) Phase transitions and (B, D) P- and S-wave velocities and density from surface to the core-mantle boundary in the Earth’s and Martian mantle. Red lines show temperature profile in the mantle. Figures for Earth are based on the mantle compositional model of [McDonough (2014)] and model geotherm from [Katsura et al. (2010)]. Figures for Mars are reprinted from T. Yoshizaki and W.F. McDonough (2020) The composition of Mars, Geochimica et Cosmochimica Acta 273, 137–162, Copyright (2020), with permission from Elsevier. Abbreviations: Brg—bridgmanite; Ca-pv—Ca-perovskite; Cpx—clinopyroxene; C2/c—high-pressure clinopyroxene; Fp—ferropericlase; Gt—garnet; Ol—olivine; Opx—orthopyroxene; Pl—plagioclase; Ring—ringwoodite; Sp—spinel; St—stishovite; Wad—wadsleyite.
Fig. 6: Interior structure of Earth and Mars. Abbreviations: Ol–olivine; Px–pyroxene; Gt–garnet; Wad–wadsleyite; Ring–ringwoodite; Brg–bridgmanite; Cpv–Ca-perovskite; Ppv–post-perovskite; Fper–ferropericlase. The Mars figure is reprinted from T. Yoshizaki and W.F. McDonough (2020) The composition of Mars, Geochimica et Cosmochimica Acta 273, 137–162, Copyright (2020), with permission from Elsevier.
Fig. 7: Composition of metallic cores of Earth (McDonough, 2014), Mars (Yoshizaki and McDonough, 2020) and NC-group iron meteorite parent bodies (Wasson and Richardson, 2001; Chabot, 2018). (A) Siderophile and chalcophile element and (B) nominally lithophile element abundances. Elemental abundances are normalized to CI chondrite composition and Ni. Elements are arranged by their 50% nebular condensation temperatures (Wood et al., 2019), while W and Re are replaced to highlight chondritic highly siderophile element abundance in planetary cores. Type IIIAB and IVA iron meteorites have fractional crystallization (i.e., magmatic) origin and are isotopically classified as NC group meteorites (Burkhardt et al., 2011; Kruijer et al., 2017a; Poole et al., 2017).
Fig. 8: Composition of metallic cores of Earth (McDonough, 2014), Mars (Yoshizaki and McDonough, 2020) and fractionally crystallized (i.e., magmatic) iron meteorites (Goldstein et al., 2009). (A) Ge versus Ni. (B) Ir versus Ni. Iron meteorites are classified into non-carbonaceous (NC: in red) and carbonaceous (CC: in blue) groups based on their Mo and W isotopic compositions (Burkhardt et al., 2011; Kruijer et al., 2017a; Poole et al., 2017). Silicate-bearing (non-magmatic) iron groups are not shown. The IIG iron meteorites are also excluded from the plot because isotopic data to classify them into NC or CC group are not available. CI chondritic ratios are from Lodders (2020).
Fig. 9: Na/Ti vs Mn/Na in chondrites (Alexander, 2019a,b), differentiated asteroids (O’Neill and Palme, 2008), Moon (Dauphas et al., 2014), Earth (this study; McDonough, 2014; Siebert et al., 2018) and Mars (Yoshizaki and McDonough, 2020). Dark gray line shows incomplete condensation from a gas of CI composition. Light gray lines correspond to evaporative loss of Na and Mn from CI and the bulk silicate Earth (BSE) compositions, respectively. See Section A.1.1 for details of model calculations.
Fig. 10: Lithophile element composition of Earth, Mars, and carbonaceous chondrites and their components. (A) Lithophile element abundance in the bulk silicate Mars (BSM), bulk silicate Earth (BSE) and bulk chondrules from carbonaceous chondrites. Elemental abundances are normalized to CI chondrite composition and Al. Elements are arranged by their 50% nebular condensation temperatures. (B) CI-normalized Al abundances in the BSE, BSM, bulk planets, bulk chondrules and bulk chondrites. Chemical composition of chondrules is from Hezel et al. (2018b) and MetBase (1994–2017). Other data sources are as in Fig. [1].
Fig. 11: (A) Accretion timescale (Myr) vs K/Th ratio of chondrites and terrestrial planets. Duration of accretion of chondritic asteroids is based on the Pb-Pb, Hf-W and Al-Mg ages of chondrules (Amelin et al., 2002; Amelin and Krot, 2007; Connelly et al., 2008, 2012; Connelly and Bizzarro, 2009; Bollard et al., 2017, 2019; Villeneuve et al., 2009; Nagashima et al., 2017; Schrader et al., 2017; Kita et al., 2013; Pape et al., 2019; Budde et al., 2016, 2018), Mn-Cr ages of asteroidal secondary alteration products (Fujiya et al., 2012; Doyle et al., 2015), and thermal modeling of asteroids (Sugiura and Fujiya, 2014). Accretion timescales of Earth and Mars are defined based on Kruijer et al. (2017b, 2020), Kleine et al. (2009), Dauphas and Pourmand (2011), Kruijer et al. (2017b), and Connelly et al. (2019). (B) Abundance of chondrules (vol%) vs K/Th value of chondrites. Error bars (shown only for carbonaceous chondrites) represent 2 standard deviations, and gray area corresponds to the 95% confidence interval of the linear regression. The abundances of chondrules in chondrites are based on McSween (1977a, 1977b), McSween (1979) and Scott and Krot (2014). The amounts of chondrules in Earth and Mars are estimated based on their K/Th values and extrapolation of the carbonaceous chondritic trend. See Section A.4 for a method to calculate the CAI-free bulk CV composition.
Fig. 12: A cartoon summarizing three models of the Earth’s accretion: (A) this study; (B) O’Neill (1991b); and (C) Wänke et al. (1984). The later models envisage Earth’s formation as composed of four stages: (1) accretion and differentiation of a proto-Earth, (2) addition of a giant impactor to the proto-Earth’s mantle, (3) loss of a sulfide matte from this mantle, and (4) late accretion of volatile-rich chondritic materials (not shown).
Fig. 13: Siderophile (red square) and chalcophile (green triangle) element abundances in the BSE, and a model BSE composition after the sulfide matte subtraction (dark red arrow) predicted by a Mars-like (i.e., size and composition) Moon-forming impactor scenario. The dark red arrows represent the model compositional range based on sulfide-silicate partition coefficients of elements (Table A.1). Highly siderophile and chalcophile elements are not shown, as their abundances in the present-day BSE can be explained by the late addition of volatile-rich materials (Step 4). Also shown are the Mars-like impactor’s contribution (red band) in the present-day Earth composition (blue band). The values for x and y axes follow the same conventions as Fig. 4.
Fig. 14: Abundances of moderately volatile elements ($X_i$) in bulk silicate Moon and mixture of the Mars-like Moon-forming impactor’s mantle and proto-BSE, plotted against mass fraction of the impactor in Moon. $X_i^{\text{bulk silicate Moon}}$ are based on Warren (2005), McDonough et al. (1992), and O’Neill (1991a). Colors of curves correspond 1% evaporation temperatures of elements from a silicate melt at log $f_{O_2} = -10$ at 1 bar (Sossi et al. 2019).
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Appendix A  Supplementary materials

A.1 Updates in planetary compositions

Here we revise compositional models of Earth (McDonough and Sun, 1995; Palme and O’Neill, 2014; McDonough, 2014) and Mars (Yoshizaki and McDonough, 2020) for some elements based on recent literature (Braukmüller et al., 2018; Alexander, 2019a,b; Wood et al., 2019; Lodders, 2020).

A.1.1 Manganese in the bulk Earth and its core

The BSE has a lower Mn/Na value than that predicted by a condensation model based on the BSE’s Na/Ti value (Fig. 9). This Mn depletion in the BSE reflects its partial incorporation into the metallic core (McDonough, 2014), which is consistent with moderately siderophile behavior of Mn under highly reduced condition in the proto-Earth (Siebert et al., 2018). Thus, the Mn abundance in the bulk Earth includes contributions from the core and the mantle.

The Mn abundance in the BSE is ~1,050 ppm (McDonough and Sun, 1995; Palme and O’Neill, 2014). McDonough (2014) estimated Mn abundance of the bulk Earth and core of 800 ppm and 300 ppm, respectively, based on a linear trend in a plot of Na/Ti vs Mn/Na among chondrites. This estimate gives core-mantle partition coefficient of Mn ($D_{Mn}^{core−mantle}$) of ~0.3, lower than that predicted by high P-T experimental studies (e.g., Siebert et al., 2018). Recently, Siebert et al. (2011) re-evaluated the bulk Earth’s bulk Mn content of 900–1300 ppm based on moderately siderophile behavior of Mn under Earth’s core formation conditions.

Here we test if the ratios of Na, Mn and Ti of Earth (McDonough and Sun, 1995; Palme and O’Neill, 2014; McDonough, 2014; Siebert et al., 2018), Mars (Yoshizaki and McDonough, 2020) and other small solar system bodies (O’Neill and Palme, 2008; Wasson and Kalsoemeyn, 1988; Braukmüller et al., 2018) reflect incomplete condensation of a solar nebular gas or post-accretionary evaporative loss of moderately volatile elements (MVE). We modeled the incomplete condensation in
the solar nebula, in which volatile depletion in condensates is strictly depending on the 50% condensation temperature of elements, following Cassen (1996) and Cassen (2001). We assume that a log of CI-normalized abundance of non-refractory elements in condensates shows a linear trend when plotted against the 50% condensation temperature of elements during a partial condensation. Thus, the composition of condensates is formulated as:

$$\log \frac{C^\text{cond}_i}{C^\text{CI}_i} = \begin{cases} -(\log T_c(\text{ref}) - \log T_c(i)) \times f & (T_c(i) < T_c(\text{ref})) \\ 1 & (T_c(i) \geq T_c(\text{ref})) \end{cases}$$ (A.1)

where $C^\text{cond}_i$ and $C^\text{CI}_i$ are abundances of an element $i$ in a condensate and CI chondrite, respectively, $T_c(i)$ and $T_c(\text{ref})$ are 50% condensation temperatures of the element $i$ and vanadium, which has the lowest condensation temperature among refractory elements (1370 K at 10 Pa; Wood et al., 2019), respectively, and $f$ is a slope of the linear volatility trend in a $\log(C^\text{cond}_i/C^\text{CI}_i)$ vs $\log T_c$ space (i.e., degree of volatile depletion in the condensate). Condensation temperature of elements from Wood et al. (2019) and CI abundance from Lodders (2020) are adopted in the calculation. The evaporative losses of elements from a high-temperature liquid were modeled using equations and thermochemical data from Sossi et al. (2019) and Chase (1998).

Our simplified calculation of incomplete condensation well reproduces a chemical composition of chondrites and Mars (Fig. 9). The bulk Earth composition (McDonough, 2014) with the updated Mn abundance (Siebert et al., 2018) is consistent with the incomplete condensation model. In contrast, compositions of differentiated asteroids (e.g., parent bodies of angrites and eucrites) and Earth’s moon are consistent with the evaporation from a CI and the BSE compositions, respectively. These observations, combined with a lack of heavy MVE isotope enrichment composition of terrestrial samples (e.g., K, Fe, Si, Mg, Zn, Cd; Humayun and Clayton, 1995; Poitrasson et al., 2004; Norman et al., 2006; Wombacher et al., 2008; Herzog et al., 2009; Paniello et al., 2012; Zambardi et al., 2013; Sossi et al., 2016, 2018), document that the bulk Earth’s MVE abundance is established by a nebular condensation, rather than post-accretionary evaporative losses. With the bulk Earth’s Na/Ti $\sim 2.2$ (McDonough, 2014), the incomplete condensation model predicts Mn abundance of 1070 ppm and 1120 ppm in the bulk Earth and its core, respectively, with the $D^\text{core-mantle}_{\text{Mn}}$ of $\sim 1.1$, which are in agreement with the values predicted by Siebert et al. (2018). Thus, we update the Mn abundances in the bulk Earth and its core with these values.

### A.1.2 Sulfur in the bulk Earth and its core

The S content in the bulk Earth is of particular interest when estimating a mass fraction of the post-impact sulfide matte. Sulfur is a moderately volatile, chalcophile element so that its abundances in the bulk Earth and core can be constrained by an abundance of lithophile element with similar volatility. Zinc abundance in the BSE has been used to constrain Earth’s S content, since Zn does not show deviation from Earth’s lithophile volatility trend and its condensation temperature is close to that of S (McDonough and Sun, 1995; Dreibus and Palme, 1996).
McDonough (2014) estimated sulfur in the BSE at 250 ± 50 ppm and a bulk Earth with 6350 ppm, based on Earth’s volatility trend and S abundance of CI chondrite of 5.40 wt%. These estimates lead to the prediction of 1.9 wt% S in Earth’s core. The authors used 50% condensation temperatures of elements from Wasson (1985), in which \( T_c \) of S and Zn (treated as lithophile) are 648 K and 660 K, respectively.

Dreibus and Palme (1996) assumed a CI-like Zn/S value for Earth given a nearly similar \( T_c \) of S and Zn (Wasson 1985), and estimated 5600 ppm S in the bulk Earth. They used a lower and higher S abundance in the BSE (130 ppm) and CI chondrite (5.9 wt%) than the values taken by McDonough (2014), which gives an estimate of \( \sim \)1.7 wt% S in the Earth’s core.

Here we revise the S content of the bulk Earth and its core based on recently-revised \( T_c \) (Wood et al., 2019) and CI abundance (Lodders, 2020). The updated \( T_c \) of S and \( T_c \) of Zn are 672 K and 704 K, respectively, with a difference of 32 K, which is larger than that of the Wasson (1985)’s value. Thus, we consider that the Zn/S value of the bulk Earth might not be chondritic if these elements are depleted in Earth based strictly on their relative volatilities. We use the S abundance of 250 ppm in the BSE (McDonough and Sun, 1995; Palme and O’Neill [2014]) and 5.36 wt% in CI chondrite (Lodders, 2020), and modeled an incomplete condensation of moderately volatile elements as we applied for Mn (Section A.1.1). With these values, we estimate S concentrations of 5900 ppm in the bulk Earth, and 1.8 wt% in the core. The revised S abundance in Earth is used to estimate the mass of post-impact sulfide matte (Section 3.3.2).

**A.1.3 Sulfur in the Martian core**

The S content in the Martian core estimated by Yoshizaki and McDonough (2020) is much lower than that estimated by Wänke and Dreibus (1994) and Taylor (2013), who constrained the S content in the Martian core by assuming that moderately volatile elements are in chondritic abundances in the bulk Mars. However, the trend of depletion of moderately volatile elements in Mars is inconsistent with assumptions in (Yoshizaki and McDonough, 2020).

Physical models of Martian interior (e.g., Khan and Connolly, 2008; Rivoldini et al., 2011; Khan et al., 2018) based on available observations (mass, density, MOI and tidal Love number \( k_2 \)) typically use the compositional model of the Martian mantle established by Wänke and Dreibus (1994). These studies generally predict a larger core (1700–1800 km in radius) with a high S concentration (15–20 wt%), which are differ from the estimates by Yoshizaki and McDonough (2020) (\( r \sim 1600 \) km: \( \leq 7 \) wt% S). These geophysical models use the Mars’ \( k_2 \) Love number to estimate its core S content. However, a deconvolution of the \( k_2 \) Love number requires an accurate knowledge of the degree of partial melting and hydration in the mantle, phase- and composition-dependent changes in elastic properties of mantle minerals, and grain size variation (e.g., Nimmo and Paul, 2013; Khan et al., 2018). Thus, we use the planetary volatility trend to estimate the Mars’ bulk and core S abundances (Yoshizaki and McDonough, 2020), following the practice used to constrain the S abundance in the Earth’s core (Dreibus and Palme, 1996; McDonough, 2014). A direct seismic
observation of Martian interior by NASA’s ongoing InSight mission will place strong constraints on the size and light element inventory of the Martian core.

Core-mantle partitioning behaviors of refractory elements, which are sensitive to the S content of the metal (Righter, 2011; Wade et al., 2012), provide further constraints on the S concentration in the Martian core. The compositional model of Mars (Yoshizaki and McDonough, 2020) predict core-mantle $D$-value of $10 \pm 5.5$, based on the depletion of W in the BSM. Figs. A.10 and A.11 show the metal-silicate partitioning behavior of W at $\Delta IW = -1.5$ for the Martian core models (Wänke and Dreibus, 1994; Taylor, 2013; Yang et al., 2015; Yoshizaki and McDonough, 2020), estimated based on a compilation of previous experimental constraints (Righter, 2011). Fig. A.10 shows that S become less siderophile as the S concentration in the metal increases. For a S-rich core model (>20 wt%; Taylor 2013), the $D$-value of W is much lower than the Martian compositional model (Fig. A.11C). Models with 10–15 wt% S in the core (Wänke and Dreibus, 1994; Yang et al., 2015) requires ~2500 K to explain the BSM’s W depletion. The S-poor core composition (S ~ 7 wt%; Yoshizaki & McDonough, 2020) is consistent with a BSM model forming at >2300 K. Thus, the predicted W abundance in the Martian core is consistent with lower S models for the Martian core. We note that the temperature conditions required to explain the distribution of W in Mars decreases by ~200 K if the log $f_{O_2}$ value decreased for 0.5 units in the $\Delta IW$ space.

Similarly, the behavior of Mo is also sensitive to S contents in a metal (Wade et al. 2012). Therefore, Mo is another useful element for constraining the S abundance in the Martian core. Unfortunately, the Mo abundance in the BSM is currently poorly constrained. Yoshizaki and McDonough (2020) found no clear correlation of Mo with other elements in Martian meteorites. Yang et al. (2015) proposed the Mo abundance in the BSM based on Mo-Ce correlation in two Martian meteorites, which were not identified by Yoshizaki and McDonough (2020). Future constraints on the Mo abundance in the Martian mantle will provide important constraints on the light element composition of the Martian core.

A.1.4 Halogens in Earth and Mars

Abundances of heavy halogens (Cl, Br and I) in the BSE is enigmatic because of their large depletion in the BSE (McDonough and Sun, 1995; Palme and O’Neill, 2014), perhaps indicative of unique behaviors of these elements during and/or after the Earth’s accretion (e.g., Kramers, 2003; Armytage et al., 2013; McDonough, 2014; Sharp and Draper, 2013; Zolotov and Mironenko, 2007; Clay et al., 2017; Jackson et al., 2018; Steenstra et al., 2020). The present bulk Earth model (McDonough, 2014) attributed this heavy halogen depletion to their incorporation into the core, and estimated core/mantle enrichment factors of 10–15 for these elements. Limited numbers of experimental studies suggest that iodine becomes siderophile during core formation (Armytage et al., 2013; Jackson et al., 2018), whereas Cl does not show such a metal-loving behavior (Sharp and Draper, 2013).

Recently, Clay et al. (2017) proposed a lower halogen abundance for chondritic meteorites than
previous estimates (e.g., Dreibus et al., 1979; Lodders, 2003). By normalizing the BSE abundance of halogens to their updated CI composition, Clay et al. (2017) showed that Br and I are plotted on the lithophile volatility trend and no need for any special processes of halogen fractionation. However, even if normalized to Clay et al. (2017)’s CI chondrite composition, the BSE’s Cl abundance still shows a depletion from the volatility trend.

The CI abundances of heavy halogens proposed by Clay et al. (2017) has been challenged by Fegley et al. (2020), who showed that these updated CI abundances fall below a curve of nuclide abundance vs mass number. Lodders (2020) critically evaluated and updated the CI composition proposed by Clay et al. (2017) and brought it in line with previous estimates of the CI abundance (Dreibus et al., 1979; Lodders, 2003).

The revised 50% condensation temperatures of the halogens (Fegley and Lodders, 2018; Wood et al., 2019) are lower than the previous values (Lodders, 2003). Consequently, using condensation temperatures from Fegley and Lodders (2018) and CI abundance from Lodders (2020), Fegley et al. (2020) showed that all the halogens plot on the Earth’s volatility trend for the lithophile elements, thereby eliminating any need for any extra halogen fractionation (Fig. A.3A). We note that a hockey-stick depletion pattern of MVE in the BSE (e.g., Braukmiller et al., 2019) is not observed when CI abundance from Lodders (2020) is used instead of that of Clay et al. (2017). In this case, the Earth’s halogen budgets (i.e., 12 ppm Cl, 0.03 ppm Br, and 0.01 ppm I in the bulk Earth) are hosted in its silicate fraction, and there is no need for their incorporation into the core.

In contrast, a separate effort by Wood et al. (2019) derived higher 50% condensation temperatures for these elements compared to Fegley and Lodders (2018), which seem to be consistent with the halogen-bearing core model (Fig. A.3B). Thus, further analytical, experimental and theoretical efforts are needed to better constrain the abundance and distribution of heavy halogens in Earth.

Yoshizaki and McDonough (2020) estimated halogen abundances in the BSM and bulk Mars based on the CI abundance of these elements from Clay et al. (2017). Using CI chondritic abundance from Lodders (2020), and assuming that the heavy halogens are hosted in the Mars’ silicate fraction, the BSM and bulk Mars heavy halogen abundances are updated as follows: 28 ppm Cl, 0.13 ppm Br, and 0.03 ppm I in the BSM; and 23 ppm Cl, 0.10 ppm Br, and 0.02 ppm I in the bulk Mars. If the Martian core contained heavy halogens, the bulk Mars’ halogen abundance might be 0.1–0.2 × CI.

A.2 Heat production in terrestrial planets

Internal heat production of Mars is calculated using the BSM abundances of heat-producing elements (HPE) $^{40}$K, $^{232}$Th, $^{235}$U and $^{238}$U (Yoshizaki and McDonough, 2020) and their standard decay constants (McDonough et al., 2020 and references therein). The Martian core contributes negligibly to the radiogenic heating of the planet as it is predicted to contain insignificant quantities
of HPE. Heat production in the Martian crust is calculated for a crust with 3740 ppm K, 700 ppb Th and 180 ppb U (Taylor and McLennan, 2009). Abundances of HPE in the Martian mantle are based on mass-balance considerations

$$X_{\text{Mm}}^i = \frac{X_{\text{Mc}}^i \times M_{\text{Mc}} - X_{\text{BSM}}^i \times M_{\text{BSM}}}{M_{\text{Mm}}} \quad (A.2)$$

where $X_j^i$ is concentration of HPE i ($^{40}$K, $^{232}$Th, $^{235}$U and $^{238}$U) in the reservoir j (Mm, Mc, and BSM are Martian mantle, Martian crust and the bulk silicate Mars, respectively) and $M_j$ is mass of reservoir j (Table 1). For $X_{\text{BSM}}$ of 360 ppm K, 68 ppb Th and 18 ppb U (Yoshizaki and McDonough, 2020) and $X_{\text{Mc}}$ from Taylor and McLennan (2009), Eq. (A.2) yields 190 ppm K, 36 ppb Th and 10 ppb U in the Martian mantle. Martian planetary Urey ratio ($U_r$) is given by

$$U_{r\text{Mars}} = \frac{H_{\text{BSM}}}{4\pi R^2 F} \quad (A.3)$$

where $H_{\text{BSM}}$ is heat production in the bulk silicate Mars, $R$ is surface area of Mars, $F_{\text{Mars}}$ is the average surface heat flow (19 ± 1 mW/m$^2$; Parro et al., 2017). Similarly, the internal heat production in Earth is calculated based on compositions of the BSE and Earth’s crust from McDonough (2014) and Rudnick and Gao (2014), respectively, $F_{\text{Earth}}$ from Jaupart et al. (2015), and mass and density of the crust from Wipperfurth et al. (2020).

### A.3 Rayleigh number of the Martian mantle

The Rayleigh number ($Ra$) for the internally-heated present-day Martian mantle is given by

$$Ra = \frac{\alpha \rho^2 g H d^5}{k \mu \kappa} \quad (A.4)$$

where $\alpha$ is thermal expansion coefficient, $\rho$ is density of the Martian mantle, $g$ is gravitational acceleration, $H$ is heat production in the present-day Martian mantle, $k$ is thermal conductivity, $\mu$ is viscosity and $\kappa$ is thermal diffusivity. We take $\alpha = 3 \times 10^5$ K$^{-1}$, $g = 3.7$ m/s$^2$, $k = 4$ W/m/K, $\mu = 10^{21} – 10^{23}$ Pa s, and $\kappa = 10^{-6}$ m$^2$/s (Turcotte and Schubert, 2014; Samuel et al., 2019) and obtain $Ra = 3 \times 10^5 – 3 \times 10^7$.

### A.4 CAI-free bulk composition of CV chondrite

Abundances of the MVE are apparently lower in CV chondrites than other carbonaceous chondrites, because of their higher abundance of refractory inclusions. Since relative abundance of refractory lithophile elements in Earth and Mars are distinct from that of CAIs and bulk CV, it is unlikely that the planetary MVE fractionation is inherited from CV CAI-like materials (Stracke et al., 2012; Dauphas and Pourmand, 2015; Barrat et al., 2016). Therefore, to evaluate the contribution of CV chondrules in the bulk CV composition, we calculated a CAI-free chemical composition of bulk CV
chondrites using mean composition of unaltered Allende CAIs ([Mason and Martin, 1977]), modal abundance of CAIs in CV ([Hezel et al., 2008; Rubin, 2011; Scott and Krot, 2014; Desch et al., 2018]), and the bulk CV composition ([Alexander, 2019a]). The calculated CAI-free bulk CV composition is shown in Figs. 11 and A.4.

A.5 Modeling the Earth’s accretion

Here we describe the details of formation model of the Earth (Section 3.3.2). The basic sequence of the Earth’s accretion is taken from Wänke et al. (1984) and O’Neill (1991a,b), with modifications:

1. the accretion of the proto-Earth (reduced and volatile-depleted) accompanied by continuous core-mantle differentiation (∼90% of Earth’s mass),
2. followed by a late-stage (e.g., 30–100 Myr after \( t_0 \); [Kleine et al., 2009]) Moon-forming giant impact event that adds the final ∼10% mass (oxidized and volatile-enriched) to Earth and forms a protolunar accretion disk,
3. subsequently, the mantle loses a Fe-Ni (±O) sulfide liquid (sulfide matte; [O’Neill, 1991b]) to the core (∼0.5% BSE mass),
4. and finally, the BSE receives the addition of (∼0.5% of the BSE mass) a late accretion component that brings the highly siderophile and chalcophile elements in chondritic proportions and highly volatile gases and fluids.

The bulk composition of the proto-Earth (Table A.2) is calculated by subtracting contributions of the Mars-like impactor and late-added materials from the bulk composition of the present-day Earth ([McDonough, 2014]). In this scenario, the bulk proto-Earth contains ≥80% of the present-day Earth’s budgets of most of the MVE (e.g., K/Th ∼ 3200; Rb/Sr ∼ 0.026; Table A.2), since a Mars-like impactor contributes only a limited amount of additional MVE (Fig. 13).

The core-mantle differentiation of proto-Earth is modeled using the calculated bulk proto-Earth composition and metal-silicate distribution coefficients of elements at high P-T conditions (∼30 GPa and ∼3000 K) and \( \log f_O^2 \sim IW - 2 \) (Table A.1).

The mass fraction of the impactor’s core that equilibrated with the Earth’s mantle (\( k \)) is poorly constrained. While a small impactor might be efficiently emulsified and completely equilibrated with the proto-Earth’s magma ocean, the behavior of Mars-sized impactor’s core remains controversial (e.g., [Rubie et al., 2003, 2015; Dahl and Stevenson, 2010; Samuel, 2012; Deguen et al., 2014]). The Hf-W and U-Pb isotopic systematics and abundances of moderately siderophile element in the BSE is consistent with 0.4 ≤ \( k \) ≤ 1 ([Jacobsen, 2005; Nimmo and Agnor, 2006; Rubie et al., 2015, 2016]). Recently, Budde et al. (2019) showed that nucleosynthetic Mo isotopic composition of the BSE is well reproduced if 20–100% of equilibrium was achieved between the impactor’s core and...
the proto-Earth’s mantle (see also Kleine et al., 2020). We found that \(0.5 \leq k \leq 1\) is consistent with the BSE abundances of most siderophile and chalcophile elements (Fig. A.7).

As a consequence of the Moon-forming event, the impactor’s mantle and core are emulsified and equilibrated within the Earth’s mantle (e.g., O’Neill, 1991b; Rubie et al., 2016), providing not only lithophile, but also siderophile and chalcophile elements to the post-impact silicate Earth (Fig. 13). The Earth’s mantle might reach S saturation, given that the addition of the impactor increases the mantle S abundance, and the S concentration at sulfide saturation decreases dramatically as the mantle cools (O’Neill, 1991b; Rubie et al., 2016). Thus, an immiscible Fe-Ni (± O) sulfide liquid (post-impact sulfide matte) precipitates through a crystallizing mantle into the core due to its high immiscibility, low wetting angle, and high density (Gaetani and Grove, 1999; Rose and Brenan, 2001).

The mass of post-impact sulfide matte is calculated by assuming that all S in the post-impact BSE was stripped by a sulfide phase, and all S in the present-day BSE is derived from the late-accreted materials (Yi et al., 2000; Rose-Weston et al., 2009; Wang and Becker, 2013). Using a metal-silicate partition coefficient of S (\(D_{\text{met-sil}}^S\)) of 100 (Rose-Weston et al., 2009; Boujibar et al., 2014), the proto-Earth’s mantle is estimated to contain ~170 ppm S. The bulk Mars-like, Moon-forming impactor has 1.2 wt% S (Yoshizaki and McDonough, 2020), so its addition increases S concentration in the post-impact Earth’s mantle to ~0.2 wt%. If all S in the post-impact BSE is segregated by a sulfide liquid with a composition of sulfides in spinel lherzolite xenoliths (37.7 wt% Fe, 21.1 wt% Ni, 0.3 wt% Co, 1.6 wt% Cu, and 38.3 wt% S; Lorand and Conquéré, 1983), the amount of the sulfide liquid is ~0.5 wt% of the present-day Earth’s silicate mantle. Fractionation of other siderophile or chalcophile elements due to the sulfide matte precipitation is modeled using the \(D_{\text{sul-sil}}^{\text{sil}}\) values listed in Table A.1. Since there are limited numbers of experimental dataset on \(D_{\text{sul-sil}}^{\text{sil}}\) values, we do not specify P-T conditions of the sulfide matte formation. The errors accompanied with the wide variation in the expected \(D_{\text{sul-sil}}^{\text{sil}}\) values are much larger than those from other parameters (e.g., planetary compositional models, P, T-dependent \(D_{\text{met-sil}}^{\text{met}}\) values), requiring additional experimental efforts in future.

The elevated abundances of highly siderophile and chalcophile elements in the Earth’s mantle indicate an addition of ~0.5% of volatile-rich materials at the final stage of Earth’s accretion (e.g., Kimura et al., 1974; Chou, 1978; Walker et al., 2015; Rubie et al., 2016; Wang and Becker, 2013; Righter et al., 2018a). The absence of meteorite class that matches relative abundances and isotopic composition of highly siderophile or chalcophile elements in the BSE keeps the nature of this late-accreted material controversial, but there is a general consensus that the late-added material had a chondritic elemental composition (Walker et al., 2002; Albarede, 2009; Alexander et al., 2012; Wang and Becker, 2013; Varas-Reus et al., 2019; Fischer-Gödde et al., 2020). In this modeling, we do not specify the type of chondritic materials added during the final stage. Note that all classes of chondrites are similarly enriched in volatiles compared to the rocky planets, and mass fraction of the late-accreted material in the Earth’s mantle is only ~0.5 %. Therefore, the choice of other type
of chondrites (e.g., enstatite or ordinary) as the late-accreted material do not affect the main results of this modeling.

The sulfide matte segregation decreases the abundances of the siderophile and chalcophile elements to the present-day BSE levels (Fig. 13). It also removes HSE like Au, Pd, Pt from the Earth’s mantle, decreasing their mantle abundance >10 times smaller than the present-day values (Rubie et al., 2016; Righter et al., 2018a). As a final step, the late addition of ~0.5 wt% of chondritic materials to the Earth’s mantle after the sulfide matte segregation increases mantle abundances of HSE and chalcogens to the present-day BSE level (Wang and Becker, 2013; Walker et al., 2015) (not shown in Figs. 13 A.6 and A.7). Thus, this accretion scenario successfully reproduces the abundances of the chalcophile and highly siderophile elements in the BSE.

A challenge to our model is the unexplained higher levels of Cu and Sn and strong depletion Sb and Ge in the BSE. These misfits can be due to our limited understanding of chalcophile behavior of these elements at the high P and T conditions. The problem with the Cu abundance is also found even in the CI-like Impactor model (O’Neill, 1991b), and it would be relaxed if the lower BSE abundance of Cu (20 ppm instead of 30 ppm; O’Neill, 1991b; McDonough, 2014) and/or lower $D_{\text{Cu}}^{\text{sul-sil}}$ under oxidizing or high-T conditions (e.g., Li and Audétat, 2015) are considered.
Table A.1: Metal-silicate and sulfide-silicate partition coefficients of elements (D) adopted in the modeling.

| Element | \(D_{\text{met} - \text{sil}}\) | Reference | \(D_{\text{sul} - \text{sil}}\) | Reference |
|---------|-------------------------------|-----------|-------------------------------|-----------|
| P       | 20                            | R10       | 10–300                        | JD86, R97 |
| S       | 100                           | R09, B14, S17 | \(\gg 1000\) | See note[b] |
| V       | 1.3                           | R11       | 0.1–2                         | GG97      |
| Cr      | 2                             | R11       | 2–4                           | KW13      |
| Mn      | 0.8                           | R11, S18  | 1–3                           | KW13      |
| Fe      | 20                            | Mass balance | 9                         | Mass balance[d] |
| Co      | 30                            | R11       | 40–100                        | KW13, P13 |
| Ni      | 40                            | R11       | 100–800                       | KW13, P13, WW17 |
| Cu      | 15                            | R11       | 100–500                       | KW13, B15 |
| Zn      | 0.5                           | Y15       | 2–5                           | KW13, P13 |
| Ge      | 200                           | Y15       | 0–5                           | KW15      |
| As      | 50                            | R17       | 1–1000                        | KP89, LA15 |
| Se      | 1000                          | R09       | 1000–2000                     | B15       |
| Mo      | 50                            | R11       | 10–1000                       | KW13, P13, LA15 |
| Ag      | 50                            | W14       | 400–1000                      | KW13, P13, B17 |
| Cd      | 2                             | R18       | 40–100                        | KW13      |
| Sn      | 50                            | R18       | 10–100                        | P13, LA15, B17 |
| Sb      | 100                           | R17       | 10–100                        | KW13, B15 |
| W       | 20                            | R11       | 1–200                         | JD86, KP89, R97 |
| Pb      | 25                            | WH10      | 5–200                         | KW13, P13, B15, B17 |

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\[a\] B14–Boujibar et al. (2014); B15–Brenan (2015); B17–Ballhaus et al. (2017); GG97–Gaetani and Grove (1997); JD86–Jones and Drake (1986); KP89–Klock and Palme (1988); KW13–Kiseeva and Wood (2013); LA15–Li and Audéat (2015); LP91–Lodders and Palme (1991); P13–Patten et al. (2013); R09–Rose-Weston et al. (2009); R97–Righter and Drake (1997); R10–Righter et al. (2010); R11–Righter (2011); R17–Righter et al. (2017); R18–Righter et al. (2018b); S17–Suer et al. (2017); S18–Siebert et al. (2018); Y15–Yang et al. (2015); W14–Wood et al. (2014); WH10–Wood and Halliday (2010); WW17–Wohlers and Wood (2017).

\[b\] Assumed that all S in the post-impact Earth’s mantle were subtracted by the sulfide matte with a present-day mantle sulfide composition (Lorand and Conquéré 1983).

\[c\] Based on mass balance between proto-Earth, Mars-like impactor, post-impact sulfide matte and present-day silicate Earth.

\[d\] Based on mass balance between post-impact Earth’s mantle and sulfide matte with a present-day mantle sulfide composition (Lorand and Conquéré 1983).
Table A.2: Composition of the bulk proto-Earth predicted by the Mars-like Moon-forming impactor model. Concentrations are in ppm (µg/g), otherwise noted.

| Element | Bulk proto-Earth | Element | Bulk proto-Earth |
|---------|------------------|---------|------------------|
| Li      | 1.1              | Nb      | 0.4              |
| Be      | 0.05             | Mo      | 2                |
| B       | 0.1              | Ag      | 0.04             |
| O (%)   | 29.0             | Sn      | 0.2              |
| F       | 9                | Sb      | 0.04             |
| Na      | 1600             | Cs      | 0.03             |
| Mg (%)  | 15.4             | Ba      | 4.5              |
| Al (%)  | 1.59             | La      | 0.44             |
| Si (%)  | 16.0             | Ce      | 1.1              |
| P       | 660              | Pr      | 0.17             |
| S       | 5200             | Nd      | 0.84             |
| K       | 180              | Sm      | 0.27             |
| Ca (%)  | 1.90             | Eu      | 0.10             |
| Sc      | 11               | Gd      | 0.37             |
| Ti      | 810              | Tb      | 0.07             |
| V       | 110              | Dy      | 0.46             |
| Cr      | 4680             | Ho      | 0.10             |
| Mn      | 920              | Er      | 0.30             |
| Fe (%)  | 32.9             | Tm      | 0.046            |
| Co      | 900              | Yb      | 0.30             |
| Ni (%)  | 1.87             | Lu      | 0.046            |
| Cu      | 60               | Hf      | 0.19             |
| Zn      | 30               | Ta      | 0.025            |
| As      | 1.7              | W       | 0.17             |
| Rb      | 0.3              | Pb      | 0.17             |
| Sr      | 13               | Th      | 0.055            |
| Y       | 2.9              | U       | 0.015            |
| Zr      | 7.1              |         |                  |
Fig. A.1: Comparison of lithophile element abundances in models of the bulk silicate Mars composition (Yoshizaki and McDonough 2020; Taylor 2013; Wänke and Dreibus 1994; Morgan and Anders 1979; Lodders and Fegley 1997). Condensation temperatures of elements are from Lodders (2003) (A) and Wood et al. (2019) (B), and CI composition is from Lodders (2020). RLE–refractory lithophile elements.
Fig. A.2: Bulk major element compositions by atomic proportions for Earth (McDonough, 2014), Mars (Yoshizaki and McDonough, 2020) and chondritic meteorites (Alexander, 2019a,b).
Fig. A.3: Abundances of moderately volatile and volatile elements in Earth (this study; McDonough, 2014), Mars (this study; Yoshizaki and McDonough, 2020) and chondritic meteorites (Alexander, 2019a). The bulk Earth model with a halogen-bearing core (McDonough, 2014) is also shown. The x axis corresponds to the 50% condensation temperature of elements at 10 Pa from Lodders (2003) and Fegley and Lodders (2018) (for halogens) in the top panel, and from Wood et al. (2019) in the bottom.
Compositions of chondrites, Earth, and Mars are from Alexander (2019a,b), McDonough (2014), and Yoshizaki and McDonough (2020), respectively. The abundances of chondrules in chondrites are taken from Scott and Krot (2014). The amounts of chondrules in Earth and Mars are estimated based on their K/Th values (Fig. 11).
Fig. A.5: Lithophile element composition of Earth, Mars and non-carbonaceous chondrites and their components. (A) Lithophile element abundance in the bulk silicate Mars (BSM), bulk silicate Earth (BSE) and bulk chondrules from non-carbonaceous chondrites. Elemental abundances are normalized to CI chondrite composition and Al. Elements are arranged by their 50% nebular condensation temperatures. Note that Cr and Mn depletion in the BSE and enstatite chondrite chondrules reflects less lithophile behavior of these elements under reduced conditions. (B) CI-normalized Al abundances in the BSE, BSM, bulk planets, bulk chondrules and bulk chondrites. Chemical composition of chondrules is from Hezel et al. (2018), Yoshizaki et al. (2018) and MetBase (1994–2017). Other data sources are as in Fig. [1].
Fig. A.6: Same as Fig. 13 but different mass fractions of the impactor (10% and 15%).
Fig. A.7: Same as Fig. 13 but different equilibrium factors ($k$) between the impactor core and the proto-Earth’s mantle.
Fig. A.8: Core/mantle enrichment factors of siderophile elements in Mars (Yoshizaki and McDonough, 2020) and metal-silicate distribution coefficients of these elements modeled at 14 GPa and 2100 K (gray), 20 GPa and 2500 K (red), and 25 GPa, 2000 K (blue) following Righter and Chabot (2011) and Yang et al. (2015). Redox conditions are fixed at log $f_{O_2} = I_W - 1.5$. Error bars on the Mars' values reflect 1 standard deviations in the Martian bulk silicate composition model (Yoshizaki and McDonough 2020).
Fig. A.9: Metal-silicate partitioning coefficients of siderophile elements calculated for the Martian core model of Yoshizaki and McDonough (2020). Redox conditions are fixed at $\log f_{O_2} = IW - 1.5$. White solid and broken lines show $P-T$ conditions that are consistent with the core-mantle distribution of these elements in Mars within 1 standard deviations (Yoshizaki and McDonough 2020).
Fig. A.10: Metal-silicate partitioning coefficients of W at (A) 15 GPa and (B) 20 GPa as a function of mole fraction of S in the metal ($X_S$), calculated using parameters from Righter (2011). The redox condition is fixed at $\Delta IW = -1.5$. The gray band shows the core-mantle partitioning coefficients ($\pm 1$ standard deviation) of W based on the Mars model of Yoshizaki and McDonough (2020). The vertical dashed gray lines show the core models of Yoshizaki and McDonough (2020) (YM20), Yang et al. (2015) (Y15), Wänke and Dreibus (1994) (WD94), and Taylor (2013) (T13).
Fig. A.11: Metal-silicate partitioning coefficients of W calculated for the Martian compositional models of (A) Yoshizaki and McDonough (2020), (B) Wänke and Dreibus (1994), (C) Taylor (2013), and (D) Yang et al. (2015). White solid and broken lines show $P$-$T$ conditions that are consistent with the core-mantle distribution of W in Mars within 1 standard deviations (Yoshizaki and McDonough 2020).
Fig. A.12: Relative contributions of radiogenic heat to Mars during accretion over the first 16 Myr of the solar system history. The compositional model for the bulk Mars is from Yoshizaki and McDonough (2020). The $\tau_{\text{accretion}}$ age of 2 Myr is adopted in the modeling (Dauphas and Pourmand, 2011).
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