The temperature and composition of the mantle sources of Martian basalts

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The temperature and composition of the mantle sources of Martian basalts

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Key Points:

- Basalts that sampled discrete mantle regions throughout Mars’s history provide information about the mantle composition and temperature
- The mantle potential temperature of primitive basalts appears constant (1400–1500 °C), yet is likely not representative of the average mantle
- Incompatible element concentrations in the mantle vary due to magma ocean crystallization, partial melting and metasomatism

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Abstract
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Plain Language Summary
Martian meteorites and rocks analyzed by rovers are witnesses of magmatic processes on Mars. In this study, we use the mantle melting model MAGMARS to determine the composition and temperature of the mantle regions from which primitive basalts have originated. Primitive basalts are closely related to mantle melts and hence record the properties of their mantle source. We find that the mantle compositions needed to explain these melts were poor in CaO and Al₂O₃. They likely represent a mantle that melted on several occasions or that crystallized from an early magma ocean. The composition of these primitive basalts indicates that some elements (Na₂O, K₂O, P₂O₅ and TiO₂) were subsequently added to the mantle source by fluids and low-degree melts. Alternatively, these elements can be explained by the trapping of melts during the evolution and progressive crystallization of the magma ocean. The temperature of the mantle sources projected to the surface conditions for easier comparison, (i.e., potential temperature) was 1400–1500 °C, regardless of the time at which these sources melted and is within the range of the most recent predictions from planetary-scale models of interior dynamics.

1 Introduction
Our knowledge of the thermal state, composition and structure of the Martian mantle is derived from a diverse and continuously expanding array of geophysical and geochemical constraints. Early measurements of the moment of inertia factor, soil compositions at the Viking landing sites, and the definitive recognition that the “SNC meteorites” are from Mars (Baird et al., 1976; Johnston & Toksöz, 1977; Bogard & Johnson, 1983), unequivocally pointed to a FeO-rich mantle (Mg/(Fe+Mg) × 100 in moles or Mg# = 75–81) compared to Earth (90). Model compositions of the “primitive mantle” were rapidly put forth (e.g., Dreibus & Wänke, 1985) and allowed to create simple models of the Martian interior structure (Longhi et al., 1992; Bertka & Fei, 1997; Elkins-Tanton et al., 2003). Additional analyses of crustal rocks by subsequent orbiting probes and rovers, the discovery of new Martian meteorites (Agee et al., 2013; Humayun et al., 2013), geodetic and seismic data from the recent InSight mission (e.g., Khan et al., 2021; Huang et al., 2022), and geodynamic modeling (e.g., Plesa et al., 2022), are now allowing to draw ever improving representations of the interior structure of Mars and its evolution through time.

Currently available compositions of the Martian mantle (e.g., Dreibus & Wänke, 1985; Lodders & Fegley, 1997; Yoshizaki & McDonough, 2020, abbreviated as DW85, LF97, YM20 and K22 hereinafter) represent average and idealized primitive compositions that are useful to derive average characteristics (density, solidus temperature, seismic wave velocity, etc.) but that probably do not represent actual regions of the mantle. The study of Martian meteorites has long shown that the mantle is highly
heterogeneous—both in terms of isotopic composition and Mg#—and suggests that a significant portion of the crust was formed very early (20–100 Myr; e.g., Borg et al., 1997; Deballé et al., 2008; Humayun et al., 2013; Nyquist et al., 2016; Kruijer et al., 2017; Bouvier et al., 2018) during (or briefly after) the crystallization of a Martian Magma Ocean (MMO). However, the major-element composition of the mantle reservoirs formed during the early differentiation of Mars is poorly constrained and model-dependent (e.g., Borg & Draper, 2003; Elkins-Tanton et al., 2005).

To derive more detailed models of the interior structure of Mars, independent constraints on the composition and temperature of discrete regions of the Martian mantle are desirable. A subset of Martian basalts, characterized by varied crystallization ages and high Mg# have been suggested to represent primitive basalts in near-equilibrium with their mantle sources and have been used to determine the $P$–$T$ conditions of their mantle source through experiments (Musselwhite et al., 2006; Monders et al., 2007; Filiberto et al., 2008; Filiberto, Dasgupta, et al., 2010; Filiberto, Musselwhite, et al., 2010) or modeling (Lee et al., 2009; Filiberto & Dasgupta, 2011, 2015; Filiberto, 2017; Bartoux et al., 2011; Balta & McSween, 2013a). Most of these basalts cannot be produced by melting the primitive mantle and are instead expected to derive from mantle sources of diverse compositions (e.g., Schmidt & McCoy, 2010; Collinet et al., 2015, Fig. 1).

Here, we use MAGMARS, a new model developed to simulate melting in the Martian mantle (Collinet et al., 2021), to re-evaluate the melting conditions and the thermal state of the mantle sources of primitive Martian basalts, which crystallized at different times and therefore represent snapshots of Mars’ thermochemical evolution. In addition, MAGMARS allows us to estimate for the first time the major-element composition of these local mantle sources. We find that the $P$–$T$ melting conditions appear to have remained relatively stable through time and that mantle sources display variable CaO/Al$_2$O$_3$, low overall abundances of incompatible elements but enrichment of alkalis, P and Ti relative to Ca and Al. We discuss the implications of these findings for the early differentiation of Mars and its long-lived magmatism.

2 Selected compositions of primitive Martian basalts

While the majority of mantle melts were modified by igneous differentiation as they ascended through the crust (Udry et al., 2018; Payré et al., 2020; Ostwald et al., 2022; Farley et al., 2022; Wiens et al., 2022), a limited number of Martian basalts bear witness to the composition and temperature of the mantle at the time of their formation (i.e., primitive basalts). To identify primitive basalts, we first make the assumption that the average Martian mantle contains olivine Mg# $\geq$ 77 (Table 1, Table S1 and Fig. 1), and would produce primary melts with a Mg# $\geq$ 54 ($K_{D,Fe-Mg}^{\text{oliv-liquid}}$ of 0.35; Filiberto & Dasgupta, 2011). A mantle of Mg# 77 is intermediate between the most commonly accepted primitive mantle compositions (Dreibus & Wänke, 1985; Yoshizaki & McDonough, 2020). Here, we only consider martian basaltic compositions with a Mg# $\geq$ 48, which could derive from primary mantle melts of Mg# $\geq$ 54 following a maximum of 10 wt.% of olivine fractionation.

The Spirit rover analyzed numerous basalts with Mg# 48–55 at Gusev crater (McSween, Wyatt, et al., 2006; Squyres et al., 2007; Ming et al., 2008) that could represent primitive basalts (Monders et al., 2007; Filiberto, Dasgupta, et al., 2010; Schmidt & McCoy, 2010). Among these, the Adirondack-class basalts are poor in K$_2$O and could derive from a residual mantle depleted in incompatible elements by prior melting events (Schmidt & McCoy, 2010; Collinet et al., 2021) while most of the basalts analyzed in the vicinity of the Columbia Hills are more enriched in alkali elements and poorer in CaO (Fig. 1). The ancient regolith breccia NWA 7034/7475/7533 (Humayun et al., 2013; Nyquist et al., 2016; Cassata et al., 2018; Bouvier et al., 2018) is also characterized by a high Mg# (54; Wittmann et al., 2015) and, despite its complex history, could approach the com-
position of a mantle melt based on trace (Humayun et al., 2013) and major element compositions (Collinet et al., 2015). We also test whether two individual clasts could be representative of primitive basalts later remelted by impacts: a vitrophyre (Udry et al., 2014) and an alkali-rich microbasalt known as “Clast VI” (Humayun et al., 2013).

Recent geophysical constraints suggest that large portions of the mantle could be more Mg-rich (Mg# = 81; Khan et al., 2022) than previously assumed (e.g., Dreibus & Wänke, 1985; Yoshizaki & McDonough, 2020), as also evidenced by the study of Martian meteorites. The most primitive depleted shergottite (Yamato 980459, nearly identical to NWA 5789; Greshake et al., 2004; Gross et al., 2011) and the most primitive enriched shergottite (LAR 06319, nearly identical to NWA 1068; Barrat et al., 2002; Peslier et al., 2010) have Mg# of 66 and 58, respectively. Y 980459 contains olivine Mg# 85–86 and is thought to represent a primary melt composition (e.g., Musselwhite et al., 2006; Matzen et al., 2022). The olivine megacrysts in LAR 06319 and NWA 1068 have Mg# ≤ 77 (Basu Sarbadhikari et al., 2009) but were initially more magnesian (Mg# 80) and were modified by Fe–Mg diffusion (Balta et al., 2013; Collinet et al., 2017). NWA 2737 is a dunitic cumulate (Mg# 79) with olivine-hosted melt inclusions. Its primary melt is taken as the reconstructed composition of the parental trapped liquid (PTL; He et al., 2013). Given the multitude of evidence of Mg-rich mantle reservoirs, we also calculated alternative primary melt compositions for the Gusev basalts and NWA 7034/7475/7533 bulk rock and basaltic clasts that would be in equilibrium with a Mg# of 81. In this case, larger amounts of olivine have to be added to the parental melt compositions (Table 1).

Table 1. List of Martian primitive basalts, fraction of olivine addition required to reach mantle-melt equilibrium, and associated inferred mantle Mg#

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<tr>
<td>Fastball [8]</td>
<td>3.7 [6]</td>
<td>0 / +13</td>
<td>77 / 81</td>
</tr>
<tr>
<td>Stars, etc. [8]</td>
<td>3.7 [6]</td>
<td>+5 / +17</td>
<td>77 / 81</td>
</tr>
<tr>
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<td>0.47 [12]</td>
<td>0 / +7</td>
<td>85 / 86</td>
</tr>
<tr>
<td>enriched shergottite LAR 06319 [13]</td>
<td>0.19 [14]</td>
<td>0 / +5</td>
<td>80 / 81</td>
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3 Methods

To constrain the mantle sources of the target basaltic compositions described above (Table 1 and S1), we first simulate the melting of various primitive mantle compositions (DW85, YM20 and K22) using MAGMARS (Collinet et al., 2021). We then adjust the mantle compositions incrementally (Mg#, TiO₂, Al₂O₃, CaO, Na₂O, K₂O, and P₂O₅ concentrations) until the liquids produced are identical to the target compositions (i.e.,
Figure 1. Comparison between the composition of Martian primitive basalts (left extremity of colored lines), their recalculated primary melts (symbols) and the melts produced by melting of the primitive mantle of Dreibus and Wänke (1985) at 1.5 (grey line) and 3.0 GPa (black line), as calculated by MAGMARS. The high SiO$_2$ content of primary melts (A) is consistent with shallow melting conditions (< 3.0 GPa). But compared to shallow DW85 melts (1.5 GPa), many primary basalts have either lower or higher CaO contents (B) and higher Na$_2$O and K$_2$O contents (C-D), and must therefore derive from mantle sources of contrasting compositions.
when the concentrations of all major and incompatible elements are within 1 wt.% relative). Next, we mathematically remove a fraction (33 to 50 wt.%) of the melt (of composition identical to the target compositions) and repeat the same procedure to identify more refractory mantle compositions that can still produce identical melts. This approach, in the absence of independent constraints on the melt fraction, leads to the identification of several possible mantle sources for each target composition. To discuss the non-uniqueness of the sources and quantify model uncertainties for the Fastball primary melt (representative example), we performed \( \sim 500000 \) MAGMARS calculations by randomly varying the parameters around their average values. This automated search identified slightly larger compositional trends compared to the manual search. However, the mantle sources identified manually were found sufficient to discuss the mantle source origin and melting temperature. It is this dataset (Table S2) that is described in the following sections.

4 Results

The compositions of the mantle sources that can produce melts identical to the target primary basalts (Table 1 and Fig. 1) are shown in Figure 2 and reported in Table S2. Each primary basalt composition can be matched by melting a series of mantle sources characterized by various concentrations of incompatible elements (\( \text{Al}_2\text{O}_3, \text{CaO}, \text{Na}_2\text{O}, \text{K}_2\text{O} \)), both isobarically and polybarically. Despite the non-uniqueness of solutions, first-order chemical differences between the sources of the different basaltic compositions can be identified. For example, the possible sources of shergottites are all notably poorer in \( \text{Al}_2\text{O}_3 \) and \( \text{Na}_2\text{O} \) than the sources of the Gusev basalts (Fig. 2a). Among the latter, the sources of the Columbia Hills basalts are characterized by high \( \text{Na}_2\text{O}, \text{K}_2\text{O}, \) and \( \text{P}_2\text{O}_5 \) concentrations (Fig. 2b–d) compared to the source of the Adirondack basalts. The source of the NWA 2737 chassignite shows the highest \( \text{K}_2\text{O}/\text{Na}_2\text{O} \) ratio. Finally, one of the sources that can match the composition of Clast VI (NWA 7533) is nearly identical to the DW85 primitive mantle.

The melt fractions required to produce the primary basalt compositions are comprised between 5 and 30 wt.%. The associated mantle potential temperatures \( (T_p) \) are between 1320 and 1520 °C (Fig. 3a and Table S2). The average pressure of melting is relatively low for all samples (1.1–2.0 GPa), and is largely constrained by the \( \text{SiO}_2 \) and \( \text{MgO} \) concentrations of the target primary melts (Fig. 1a). If a Mg# of 81 (K22) is assumed instead of 77 for NWA 7034/7475/7533 and Gusev basalts, then the primary basalts would contain a larger olivine component and the mantle \( T_p \) (1390–1570 °C) and average pressure of melting (1.9–3.0 GPa) would both be higher (Fig. 3).

5 Discussion

5.1 Thermal state of the Martian mantle

Compared to the \( T_p \) estimates of Filiberto (2017), and using the same starting assumptions (mantle of Mg# 77 and batch melting), we find that Gusev crater basalts are derived from slightly cooler mantle sources on average, with \( T_p \) of \( \sim 1400 \) °C (vs. \( \sim 1450 \) °C), but that the ranges of possible \( T_p \) largely overlap (1360–1460 vs. 1390–1550 °C, respectively). Allowing for a higher Mg# of the mantle sources (77–81), we find that Gusev basalts and all (pre-)Noachian to Hesperian samples point to a \( T_p \) of 1340–1520 °C (Fig. 3a).

We calculate a \( T_p \) of 1420–1430 °C for the primary melt composition reconstructed from NWA 2737 melt inclusions (He et al., 2013), assumed to be parental to the middle-Amazonian nakhlites and chassignites (1.34 Ga; Udry & Day, 2018). However, the mantle source could have been metasomatized (Day et al., 2018, also see section 5.2) and could
Figure 2. Incompatible element concentrations of the mantle sources of primary basalts (symbols) compared to residual model Martian mantles (lines). Each line represents the trajectory of residues produced by progressive melting of a primitive mantle composition (apex) at 1.5 (grey) and 3.0 GPa (black). For the DW85 model (solid lines), tick marks indicate the composition of residues after specific degrees of melting (in wt.%). All mantle source compositions are relatively poor in CaO and Al₂O₃ compared to the primitive mantle and are characterized by variable CaO/Al₂O₃ (A). The source of NWA 7533 clast VI is similar to a primitive mantle (DW85) and the source of the Adirondack basalts resemble a residual mantle following prior partial melting in all compositional spaces (A–D). All other sources are too rich in alkali elements—Na₂O (B) and especially K₂O (C)—and other incompatible elements, such as P₂O₅ (D), to derive from model Martian mantles by partial melting alone and other processes must be considered (see section 5.2).
have contained up to 250 ppm of water (McCubbin et al., 2016), which would translate into a lower \( T_p \) of 1380 °C (Katz et al., 2003).

The more recent olivine-phyric shergottites (160-500 Ma; Moser et al., 2013; Wu et al., 2021; McFarlane & Spray, 2022) are picritic basalts that have been linked to plumes with a \( T_p \) of at least 1480–1550 °C (e.g., Musselwhite et al., 2006; Filiberto & Dasgupta, 2015). The \( T_p \) of MAGMARS simulations (1470–1520 °C) are within error of these previous constraints if batch melting is assumed, and slightly lower in the polybaric case (1440–1450 °C). The presence of water in the source of shergottites could in principle lower the minimum \( T_p \) and has been suggested to account for their relatively high \( \text{SiO}_2 \) concentrations (Balta & McSween, 2013b). However, the small initial water concentration of the source (14–73 ppm; McCubbin et al., 2016) and the fact that the \( \text{SiO}_2 \) concentration of shergottite melts can be reproduced with MAGMARS under nominally anhydrous conditions preclude a significant effect of water.

Finally, we re-calculate using MAGMARS the \( T_p \) and pressures of melting of the bulk volcanic provinces of Baratoux et al. (2011), as constrained by the Gamma Ray Spectrometer (GRS) on board NASA’s Mars Odyssey spacecraft. Baratoux et al. (2011) used pMELTS in their analysis, which has since been shown to overestimate FeO and underestimate \( \text{SiO}_2 \) concentrations by up to 8 wt.\% (Collinet et al., 2021), significantly more than anticipated by El Maarry et al. (2009). For Hesperian provinces, while the ranges of \( T_p \) are similar (1390–1460 vs. 1370–1420 °C previously), MAGMARS predicts a slightly higher pressure of melting (1.6–2.3 vs. 1.3–1.6 GPa). However, we find that only Ascraeus and Elysium Mons (out of the 6 Amazonian volcanic provinces) can be matched with a DW85 mantle composition using MAGMARS (Table S3). The composition of the other 4 provinces can either not be reproduced at all (Arsia and Pavonis Mons) or only with an extremely small melt fraction of <2 wt.\% (Olympus Mons and Alba Patera). With a YM20 composition (Mg# of 79, 81 after 15 wt.\% of melting), a higher \( T_p \) of 1520–1660 °C and higher pressures of melting (2.3–3.5 GPa) are necessary to match the Hesperian volcanic provinces. A higher Mg# mantle also allows to reproduce the composition of a greater number of Amazonian volcanic provinces (5, all but Arsia Mons) with \( T_p \) of 1380–1460°C and pressures of 2.8–3.1 GPa.

The lack of temperature and pressure trends over time displayed by this set of constraints renders it impossible to calculate rates of secular cooling or lithosphere thickening (Fig. 3a,b). This could be due to the limited number of primitive basalts available that might not be representative of the average mantle. To test this possibility, we compare the mantle temperature estimates derived from MAGMARS to the results of a global convection model incorporating the most recent interior structure constraints from InSight (Plesa et al., 2022). The maximum temperature (and minimum pressure) at which the mantle is melting decreases with time (i.e. secular cooling). However, at any given time, melt is produced from regions of the mantle with highly variable \( T_p \), which encompass the \( T_p \) of the mantle sources estimated in this study. The Gusev basalts are the only primitive basalts whose location is known with certainty. Additionally, NWA 7034 and the depleted shergottites have recently been suggested to have originated from Karratha and 09-000015 craters, respectively (Lagain et al., 2021, 2022). Under all three locations and at the appropriate—and highly contrasting—crystallization ages, the \( T_p \) of the mantle sources would have been nearly identical and in the range 1525–1562 °C (Fig. 3c,d and S3). This confirms that despite the overall decrease in mantle temperature with time, a limited basaltic sample suite can record near-constant mantle temperature. The thermochemical evolution model predicts that the mantle temperature should first increase due to the decay of radioactive elements and peak at the Noachian/Hesperian transition before slowly decreasing (e.g., Plesa et al., 2022). This thermal maximum is not recorded by the 3.7 billion years old Gusev basalts but seems consistent with our re-interpretation of the \( T_p \) of Hesperian volcanic provinces (1520–1660 °C; Baratoux et al.,...
2011), assuming that the average mantle is relatively MgO-rich (Mg# of 79; Yoshizaki & McDonough, 2020).

Perhaps the main discrepancy between the thermochemical evolution model and the MAGMARS constraints is the shallow depth of melting that we estimate for the source of shergottites, which is predicted to be well within the lithospheric mantle (Plesa et al., 2022). Filiberto (2017) noted that if a larger amount of olivine fractionation had taken place, the primary melts of shergottites could have been in equilibrium with the convecting mantle at 3–5 GPa. While this pressure of melting is more consistent with the thick lithosphere of the late Amazonian (Fig. S3), such melt compositions would require a high \( T_p \) of 1710 ± 73 °C, which exceeds significantly the maximum \( T_p \) achievable by thermal evolution models at that time (Fig. 3d). Therefore, we consider it more likely that the \( T_p \) of the sources was low (1470–1520 °C) and that the pressure of melting derived from MAGMARS simulations (1.6 ± 0.5 GPa) does not represent the average pressure of melting but simply the final pressure of equilibration with the mantle. If shergottites formed in the Tharsis region (e.g., Lagain et al., 2021), deeply-sourced primary melts could have re-equilibrated with a warm lithospheric mantle, locally heated by magmas, at the base of the crust (110–130 km; Wieczorek et al., 2022).

5.2 Origin of the mantle sources and their variable concentrations of incompatible elements

The mantle source of Clast VI (NWA 7533) could be nearly identical to the primitive mantle (Fig. 2), as previously suggested based on rare-earth element (REE) modeling (Humayun et al., 2013). All other mantle sources are depleted in CaO and Al\(_2\)O\(_3\) relative to the various primitive mantle compositions proposed in literature (DW85, LF97, YM20 and K22). One possibility is that these mantle sources represent melting residues from which 10–20 wt.% melt had been removed prior to producing the melts that eventually formed the primitive basalts used in this study (Fig. 2a). However, the concentrations of alkalis and other incompatible elements (e.g., TiO\(_2\), P\(_2\)O\(_5\)) are, in most cases, too high at a given Al\(_2\)O\(_3\) concentration, regardless of the style (batch vs. fractional) and pressure of melting (Fig. 2b–d). Only the Adirondack basalts are consistent in detail with the simple re-melting of a mantle residue, following ±10 wt.% prior melting of a primitive mantle (see also Collinet et al., 2021). Other processes must be invoked to explain the chemical variability of the remaining mantle sources.

The Columbia Hills basalts are often assumed to be related to the Adirondack basalts, as both groups were analyzed by Spirit at Gusev crater. Compared to the Adirondack basalts, they are rich in alkali elements as well as other incompatible elements (TiO\(_2\), P\(_2\)O\(_5\)) and poor in CaO and Al\(_2\)O\(_3\) (Fig. 1). McSween, Ruff, et al. (2006) suggested that the Columbia Hills basalts could have derived from melts similar to the Adirondack basalts by fractional crystallization. The higher incompatible element concentrations (e.g., K, P, Ti) of the Columbia Hills basalts have also been suggested to result from the contamination of Adirondack-like primitive melts by a crustal component (Schmidt & McCoy, 2010). However, crustal assimilation and fractional crystallization (AFC) of basaltic melts should lower markedly the MgO concentrations (and Mg#; Ostwald et al., 2022). As the Mg# of the Columbia Hills and Adirondack basalts are similar, most workers now regard them as two sets of near-primary melts (Schmidt & McCoy, 2010; Filiberto & Dasgupta, 2011; Collinet et al., 2015). Schmidt and McCoy (2010) proposed that the high K\(_2\)O content of the Columbia Hills basalts could be accounted for by melting a fertile mantle source with a higher K\(_2\)O content compared to the Dreibus and Wanke (1985) composition. According to their model, the Adirondack basalts would be slightly younger and produced by re-melting the same region of the mantle. However, the similarly low CaO and Al\(_2\)O\(_3\) concentrations of their sources (Fig. 2a) suggest that both the Adirondack and Columbia Hills basalts were derived from depleted mantles, affected by 10–20 wt.% prior melting at ∼3.0 GPa. Metasomatism has been invoked to reconcile the high
Figure 3. Temporal evolution of $T_p$ (A) and the average pressure and depth of melting or conditions of mantle–melt re-equilibration (B). The rectangles represent the sources of the basaltic compositions listed in Table 1. The rounded fields are the sources of the GRS volcanic provinces of Baratoux et al. (2011), re-calculated with MAGMARS. The black lines represent the evolution of the potential temperatures and pressures of the part of the mantle that is affected by partial melting in the thick-crust geodynamical model of Plesa et al. (2022). The minimum pressure of melting (dashed line in B) can be interpreted as the thinnest thermal lithosphere observed anywhere on the planet. Panels C and D represent regional variations in $T_p$ for this geodynamical model at the time of Gusev basalt (C) and depleted shergottites (D) crystallization. At their possible source locations (white stars, see text for references), the $T_p$ are nearly identical: 1562 vs. 1525 °C (see also Fig. S3).
water and incompatible element concentrations of nakhlites-chassignites with their Sr-Nd isotopic compositions indicative of ancient depleted sources (Goodrich et al., 2013; McCubbin et al., 2013; Day et al., 2018) and could also help explain the high K$_2$O concentrations analyzed in numerous rocks from Gale crater (e.g., Schmidt et al., 2014). Similarly, we posit that the relative enrichment of incompatible elements in the Columbia Hills basalts (alkali elements as well as elements like P and Ti that are less mobile in fluids) could be explained by the secondary addition of low-degree melts to a Adirondack-like mantle source. The highest possible K$_2$O concentrations that we calculate for the Columbia Hills mantle sources are in the range 0.13–0.15 wt.%. This is much smaller than the percent level K$_2$O concentrations of highly metasomatized and phlogopite-bearing terrestrial peridotites (e.g., Condamine & Médard, 2014) but similar to other intraplate peridotites containing no hydrous phases (e.g., Smith et al., 1993). The source of the Columbia Hills basalts was likely affected by low degrees of cryptic metasomatism and was thus not significantly hydrated.

The isotopic systematics of Martian meteorites suggest the existence of a magma ocean that crystallized early in Mars’ history (e.g., Elkins-Tanton et al., 2005; Debaille et al., 2008; Kruijver et al., 2017; Bouvier et al., 2018). Some of the resulting heterogeneity was never erased by convection and ancient mantle sources were affected by partial melting and formed the shergottites as recently as 170 million years ago (Moser et al., 2013; Wu et al., 2021; McFarlane & Spray, 2022). The major and incompatible element concentrations of the sources of shergottites must in part reflect the processes of magma ocean crystallization. For example, the superchondritic CaO/Al$_2$O$_3$ ratio of shergottites has been suggested to result from the fractionation of majorite in the deep mantle (Borg & Draper, 2003). Here, we find that the sources of shergottites had mildly superchondritic CaO/Al$_2$O$_3$ ratios that could have appeared at low pressure, following 15–20 wt.% melting of the primitive mantle (Fig. 2a). A 20 wt.% depletion from a primitive mantle is also sufficient to decrease the incompatible element concentrations to levels identical to those of the source of depleted shergottites (Fig. 2b–d). In this case, however, the melting residue only reaches a Mg# of 77 (when starting from a DW85 mantle) to 81 (YM20), following 20 wt.% of melting. The much higher Mg# of the source of Y 980659 (85–86) remains easier to explain if it formed as a magma ocean cumulate (e.g., Borg & Draper, 2003; Elkins-Tanton et al., 2005). The enriched shergottites have higher concentrations of incompatible elements. Their composition in radiogenic isotopes indicates that the enriched signature is most likely derived from evolved residual melts that were trapped in mantle cumulates during the crystallization of an early MMO, rather than from crustal assimilation (e.g., Borg & Draper, 2003; Symes et al., 2008; Debaille et al., 2008; Brandon et al., 2012; Ferdous et al., 2017; Armytage et al., 2018). This could also explain the slightly higher concentrations of minor incompatible elements that we calculate for the source of enriched shergottites (Fig. 2).

6 Conclusions

The mantle temperature of the sources that gave rise to known primitive basalts appears to have remained relatively stable through time ($T_p$ of 1400–1500 °C). This could be due to a sampling bias. The higher mantle $T_p$ (∼1600 °C) of the Hesperian volcanic provinces (Baratoux et al., 2011), recalculated with MAGMARS and assuming a mantle with Mg# of 79 or higher (Yoshizaki & McDonough, 2020; Khan et al., 2022), hints at a significant secular cooling (>100 °C) as expected from thermochemical evolution models (Plesa et al., 2022). The shergottite melts were likely produced at pressures greater than 3 GPa but re-equilibrated with the lithospheric mantle at 1–2 GPa, for example at the base of the thick Tharsis crust.

With the exception of the source of NWA 7034 and paired rocks, the mantle sources of known Martian basalts were poorer in Al$_2$O$_3$ and CaO compared to primitive mantle compositions (e.g., Dreibus & Wänke, 1985; Yoshizaki & McDonough, 2020). The com-
positions of the sources of Gusev crater basalts that we calculate do not explicitly re-
quire a magma ocean stage and could represent simple depleted mantle reservoirs affected
by 10–20 wt.% prior melting (Adirondack basalts) or depleted mantle reservoirs re-fertilized
by fluids and low-degree silicate melts (Columbia Hills basalts). On the other hand, the
major element composition of the source of depleted shergottites cannot be easily ex-
plained by partial melting alone and suggest, along with their Sr-Nd-Hf isotope system-
atics, that they formed as mantle cumulates during the crystallization of the MMO. The
sources of enriched shergottites are consistent with trapping a more evolved residual melt.
It is also possible that the relative enrichment of the minor incompatible elements (Na,
K, Ti and P) of the Columbia basalts is a vestige of magma ocean processes and does
not result from metasomatism. But regardless of its origin, this relative enrichment is
limited, with concentrations of Na$_2$O, P$_2$O$_5$, and TiO$_2$ not exceeding the range displayed
by primitive mantle compositions.

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Open Research
The data used for the discussion and figures is summarized in the supplementary
material (Table S1-S3) and available in full at https://doi.org/10.5281/zenodo.7691390
(Collinet et al., 2023)

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Figure 2.
Figure 3.
The image presents various graphs and maps related to thermal and geological processes in the context of planetary science. The graphs illustrate temperature-pressure relationships over time, with specific markers indicating possible source locations for craters. The maps depict temperature distributions at different geological times, indicating potential mantle temperature at 450 km. The data is sourced from the research of Plesa et al. (2022) and Baratoux et al. (2011).
The temperature and composition of the mantle sources of Martian basalts

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Additional Supporting Information (Files uploaded separately)

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2. MAGMARS results used in figure 3 to match the GRS volcanic provinces of Baratoux et al. (2011) (data repository)

3. MAGMARS scripts and files used to produce figure S1–S2 (data repository)

data repository: https://doi.org/10.5281/zenodo.7691390

S1 Non-uniqueness and model uncertainties

MAGMARS uncertainties are discussed in detail in section 3.2 of Collinet et al. (2021) and propagated by combining analytical, experimental, and model uncertainties. They are provided as fixed average uncertainties for the composition of melts produced by melting any mantle composition, at any pressure and temperature: ± 1.3 wt.% SiO₂, 0.21 wt.% TiO₂, 0.38 wt.% Al₂O₃, 0.11 wt.% Cr₂O₃, 0.85 wt.% FeO, 0.06 wt.% MnO, 0.75 wt.% MgO, 0.61 wt.% CaO, 0.28 wt.% Na₂O, 0.12 wt.% K₂O, 0.12 wt.% P₂O₅, for lherzolite melting.

To quantify the uncertainties on the mantle composition and melting conditions that can produce a specific primary basalt, we run a large number of simulations while varying systematically the parameter space (black contours in Figure S1). We retain only the simulations that produce a melt identical to the basalt Fastball (representative example) within the model uncertainties stated above (blue circles). Figure S1 can be compared to Figure 2 and shows that despite the substantial uncertainties associated with the method, the conclusions of the study remain unchanged.
Another large set of simulations is filtered assuming that the model uncertainties are smaller than reported in Collinet et al. (2021) (Figure S2). While it is possible that the MAGMARS model uncertainties are overly conservative, the goal here is simply to isolate model uncertainties and non-uniqueness. Even assuming a small model uncertainty, there is a large array of possible mantle sources and melting conditions that can produce nearly identical basaltic melts (the composition of Fastball in this case).

For a given basaltic melt (e.g., Fastball), the most refractory mantle sources are associated with the lowest melt fraction and lowest mantle temperatures due to the smaller release of latent heat of melting (Fig. S1a,b). To produce the same average basaltic liquid by polybaric melting as by isobaric melting, melting must start deeper and extend to a shallower region of the mantle, where the solidus temperature is low. The $T_p$ is therefore lower compared to the isobaric case (Table S2, Fig. S1a, S2a, also see Fig. 2 in Collinet et al., 2021).

References


Figure S1: MAGMARS simulations able to reproduce the composition of Fastball (blue circle) assuming the average model uncertainty of Collinet et al. (2021): ± 1.3 wt.% SiO$_2$, 0.21 wt.% TiO$_2$, 0.38 wt.% Al$_2$O$_3$, 0.11 wt.% Cr$_2$O$_3$, 0.85 wt.% FeO, 0.06 wt.% MnO, 0.75 wt.% MgO, 0.61 wt.% CaO, 0.28 wt.% Na$_2$O, 0.12 wt.% K$_2$O, 0.12 wt.% P$_2$O$_5$. The black envelope represents the conditions sampled by 105000 MAGMARS simulations.
Figure S2: MAGMARS simulations able to reproduce the composition of Fastball assuming a low uncertainty of ± 0.5 wt.% SiO₂, 0.05 wt.% TiO₂, 0.15 wt.% Al₂O₃, 0.11 Cr₂O₃, 0.25 wt.% FeO, 0.06 MnO, 0.25 wt.% MgO, 0.15 wt.% CaO, 0.07 wt.% Na₂O, 0.03 wt.% K₂O, 0.05 wt.% P₂O₅ (red circles). The open and closed blue triangles represent the sample sources reported in Figure 2 and Table S2. The black envelope represents the conditions sampled by 375000 MAGMARS simulations.
Figure S3: Temperature profiles of the mantle below Gusev crater, and the possible location of the sources of depleted shergottites (09-000015, Tooting; Lagain et al. (2021)) and NWA 7034 (Karratha; Lagain et al. (2022)) at the time of their crystallization, from the thick-crust model of Plesa et al. (2022). The mantle below Tooting crater does not reach the solidus of Ruedas and Breuer (2017) (grey line). Below the other 3 craters, the solidus is crossed at ~5 GPa and the mantle potential temperatures are nearly identical (filled circles on the upper temperature axis): 1556 °C (Karratha), 1562 °C (Gusev) and 1525 °C (09-000015).