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The SNC meteorites: basaltic igneous processes on Mars

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Abstract

A group of 31 meteorites (SNC group) was derived from Mars as a product of 4 – 7 ejection events, probably from Tharsis and Elysium-Amazonis. The SNCs either have basaltic mineralogy or some are ultramafic cumulates crystallised from basaltic melts. The SNCs can be classified both petrographically and geochemically. We classify the shergottite SNC meteorites on the basis of their LREE-depletion into Highly Depleted, Moderately Depleted and Slightly Depleted. The Slightly Depleted samples (which are mainly but not exclusively aphyric basalts) show high log₁₀ fO₂ values (QFM -1.0). Highly Depleted samples - which are mainly olivine-phyric basalts - have low log₁₀ fO₂ values (QFM -3.5). On the basis of mixing calculations between La/Lu and ⁸⁷Sr/⁸⁶Sr we favour models linking the correlation between LREE abundances and log₁₀ fO₂ to mantle heterogeneity rather than contamination by oxidised, LREE-rich crustal fluids. SNC chemistry in general reflects the Fe-rich mantle of Mars (x2 FeO that of the Earth), the late accretion of chondritic material into the mantle, and possibly the presence of a plagioclase-rich magma ocean which acted to variably deplete the mantle in Al. The high FeO contents of the SNC melts are associated with high melt densities (allowing the ponding of large magma bodies) and low viscosities, both of which are consistent with the large scale of many observed martian lava flows.
Of the approximately 30000 currently known meteorites, thirty two originated on Mars. The martian meteorites are also called the SNC group after 3 of its members: Shergotty, Nakhla, Chassigny. They are known to form a distinct group of meteorites from a single parent body on the basis of their relatively differentiated mineralogy and chemical compositions; oxygen isotope compositions which are related to each other by mass fractionation e.g. Clayton and Mayeda (1996); high oxidation state for meteorites log\(\text{f}_2\text{O}\) between QFM and IW e.g. Herd 2003) and notably large range of crystallization ages (165 My to 4.5 Ga, Nyquist et al. 2001a).

It is the crystallization ages that first led to wide acceptance that this meteorite group was derived from a large, slowly cooled planet – Mars (e.g. McSween et al. 1979; Wood and Ashwal, 1981). However, suggestions that some meteorites might be derived from Mars can be traced back further (e.g. Wänke, 1968). Papanastassiou and Wasserburg (1974) noted that one of the SNC meteorites (Nakhla) must have been derived from an (unspecified) planetary object with some affinities to the Earth that had undergone differentiation after 3.6 Ga. The definitive link to Mars was made through comparing the composition of gases within the shock-melted glass of shergottites to the composition of the martian atmosphere determined by the Viking Landers (Bogard and Johnson, 1983).

The SNCs have a range of basaltic and ultramafic mineral assemblages (Table 1) and we show a range of new and published mineralogical and chemical data to characterise them. Sixteen of them are basaltic shergottites and 8 of those have a subclassification as olivine-phyric. There are 6 peridotitic shergottites or ‘lherzolitic shergottites’, containing less plagioclase (maskelynite) than the basaltic shergottites. The nakhlites are a group of 6 olivine-clinopyroxenites and 1 pyroxenite; Chassigny is a dunite and ALH84001 is an orthopyroxenite. We show that the shergottites can be also be subclassified geochemically on the basis of their degree of LREE-depletion. We use new trace element data to model cumulate processes in the nakhlite magma chamber. Variable patterns of depletion in incompatible elements are a feature of the SNC source melts and can be correlated with conditions of oxygen fugacity in mantle or crustal source regions. Whether the geochemical diversity amongst the SNCs is due to mantle heterogeneity related to early magma ocean crystallization, or reflects mixing between mantle-derived magmas and crustal components is one of the issues we address in this paper. We present a new model to demonstrate the likelihood of a plagioclase-flotation magma ocean forming during Mars’ history and compare it to the lunar example.
Spectroscopic measurements (e.g. Bandfield et al. 2000) show the basaltic composition of much of the martian surface, in addition we discuss whether γ-ray, TES spectroscopic data and geochemical evidence from the 1997 Pathfinder lander results suggest more differentiated basaltic-andesites or andesites in parts of the northern lowlands as well. Regions within the northern lowlands are the most likely source of the majority of the SNCs because they have young crystallisation ages equivalent to the young ages of the northern lowland terrains inferred from crater counting. The in-situ measurements of the various landers suggest that basaltic material with some more differentiated basaltic-andesites and salts dominates the uppermost surface of Mars. The SNC meteorite compositions and rock analyses from the 2004 Spirit and Opportunity landers reflect the basalt component and as we show in this paper they are related to basaltic volcanism or cumulate processes. Siderophile element abundances in SNCs are utilised to demonstrate the effects of late chondritic accretion onto the early Mars.
Methods

Quantitative mineral analyses (EPMA) of shergottites and nakhlites were made with the Cameca SX50 at the Department of Mineralogy, Natural History Museum, UK. Accelerating voltages used were 15-20 kV with 15 nA specimen current. The X-ray maps of Governador Valadares were gathered on the Cameca SX50 at 60 nA specimen current and 20 kV. The maps obtained were then processed to normalise the data and provide colour representations across the main range of elemental concentrations. We determined trace element abundances on nakhlite SNC meteorites using a quadrupole-based laser ablation ICP-MS using NewWave Research (Fremont, USA) UP213 laser ablation system coupled to a Thermo Elemental (Winsford, UK) PQ3 ICP-MS with enhanced sensitivity S-option interface at the Natural History Museum, UK. Beam spot diameter within the rasters was 60-85 µm. For calibration we used the National Institute of Science and Technology (NIST) standard reference material NIST SRM612. Jeffries (2001) provides an in-depth discussion of the laser ablation ICP-MS technique. REE contents were determined by normalisation to Ca or Si contents previously determined by EPMA. Whole sample REE contents of Nakhla and Y000593 were determined using the ICP-MS in solution mode. The Y000593 and Nakhla (BM1913, 26) sample sizes were 100 mg. The samples were put into solution within platinum crucibles using 1ml HNO₃ + 1ml HCLO₄ + 5ml HF, evaporated to dryness and then re-dissolved in 2ml HNO₃.

Bulk analyses of the DaG476 and SaU005 shergottites and the Y000593 and Y000749 paired nakhlites were obtained at UCLA using mainly instrumental neutron activation analysis (INAA: procedure of Warren et al. 1999), augmented for major elements by EPMA of fused beads (procedure of Warren and Wasson, 1979).
The SNCs: Petrology

The basaltic shergottites have a clearly basaltic set of mineral assemblages (augite, pigeonite, plagioclase ± olivine, Table 1). The other, ultramafic SNCs are cumulates related to basaltic magmas. The SNCs are characterised by moderate to strong shock alteration i.e. 15 – 45 GPa (Nyquist et al. 2001a). This is manifested by fractures in the nakhlites and ALH84001, conversion of plagioclase to maskelynite (which is generally considered to be diaplectic glass i.e. formed in a solid state transformation) in the shergottites and ALH84001. The peridotites have undergone the highest degree of shock and post-shock heating up to ~600°C. Another feature of the SNCs related to shock alteration is glassy impact melts e.g. the lithology C veins of EET79001.

Two-pyroxene geothermometry, exsolution lamellae in shergottite and nakhlite pyroxenes and the compositions of co-existing Ti-magnetite and ilmenite all show that varying degrees of subsolidus equilibration have also taken place within many of the SNCs. For instance, ilmenite-Ti magnetite pairs within Nakhla suggest limited equilibration at 740°C (Reid and Bunch, 1975). Two-pyroxene geothermometry for the ALH84001 orthopyroxenite suggests equilibration at 875°C (Treiman, 1995).

The SNC meteorites – in particular the nakhlites and ALH84001 – also contain secondary mineral assemblages (clays, Fe-rich carbonate) which on the basis of preterrestrial fractures and truncation by fusion crust are known to be martian. These mineral assemblages contain a record of atmosphere-fluid-rock interaction and are described in Bridges et al. (2001).

Four of the SNCs (Nakhla, Chassigny, Shergotty and Zagami) are falls, most of the remainder are Antarctic (EET, QUE, Y, ALH, GRV) and desert finds (NWA, Dhofar, DaG, SaU). Three of the SNCs have uncertain origins: Los Angeles, Governador Valadares and Lafayette. These latter meteorites were identified in a dealer’s collection in Los Angeles, a university collection in Indiana, USA (Lafayette) and the city of Governador Valadares, Brazil. The desert finds show varying degrees of terrestrial weathering such as the growth of calcite.

Basaltic Shergottites

The greatest number of martian meteorites are basaltic shergottites (Tables 1, 2). They contain two clinopyroxenes (augite and pigeonite), typically zoned towards Fe-rich rims (Fig. 1). Proportions of pyroxene vary from 70 vol. % in Shergott to 44 % in QUE94201 and 43% in Los Angeles (Stolper
and McSween, 1979; Rubin et al. 2000; Warren et al. 2004) with the remainder mainly composed of altered plagioclase. The mineral modal abundances of the basaltic shergottites are also summarised in more detail by Goodrich (2003) as follows: pigeonite is 36-45%, augite 10-34%, plagioclase (maskelynite) is 22-47% with a range of bulk Mg# from 23 (Los Angeles) to 52 (Zagami). Hydrous amphiboles have been tentatively identified within melt inclusions (e.g. Floran et al. 1978; Treiman, 1985; Johnson et al. 1991). These are calculated, on the basis of the amphibole stability, to have crystallised from melts that contained up to 1.8 wt% H₂O (Johnson et al. 1991; McSween & Harvey, 1993). Watson et al. (1994) used ion probe analyses of D and H in the amphiboles to show that they actually now contain less than one tenth of such H₂O contents. They also suggested that postcrystallization D enrichment of initially D-poor phases by martian crustal fluids with near atmospheric D/H had occurred. It is possible that magmatic H₂O contents were diminished by shock and the initial values in the melts remain uncertain.

The basaltic shergottites are not all homogenous samples: Zagami is composed of three different lithologies, which although all basaltic shergottite, vary in terms of pyroxene grain size (0.2 - ~2mm) and proportions of glassy melt (McCoy et al. 1992). EET79001 is composed of a basaltic lithology B, an olivine-phyric lithology A and a lesser basaltic glassy lithology C in veins and pockets. The latter containing trapped martian atmospheric gases (Bogard and Johnson, 1983).

Plagioclase has crystallised after the pyroxenes in all shergottites and has been altered to maskelynite through shock. Ti-magnetite, ilmenite, apatite, pyrrhotite, glass and symplectites are also present within interstitial areas.

The range of pyroxene compositions is shown in Fig. 2 (and see mineral compositions of SNCs in Table 3). There is a large overall range: En₁₋₇₁Wo₀₋₃₀Fs₁₋₉₄, with Los Angeles containing the most Fe-rich pyroxenes. The other basaltic shergottites show more restricted ranges of pyroxene compositions, for instance Shergotty augites and pigeonites are En₃₀₋₇₁Wo₁₁₋₃₄Fs₁₉₋₄₇. Indirect evidence for extreme Fe-enrichment associated with pyroxene crystallization is shown by the presence of symplectites in some of the basaltic shergottites e.g. Shergotty, Zagami, Los Angeles, QUE94201 (Aramovich 2002). The symplectites are believed to have formed through replacement of ferrosilite-rich pyroxene with compositions in the ‘forbidden’ zone of the pyroxene quadrilateral during cooling above 1000°C for over 3 days (Lindsley et al. 1972). The symplectites consist of ~1-10 μm-sized fayalite + SiO₂
polymorphs ± hedenbergite and merrillite Ca$_4$(MgFe$^{2+}$)$_7$(PO$_4$)$_7$ grains. The merrillite took up Mg, forcing the crystallising pyroxene to extreme, unstable Fe-rich compositions.

Müller (1993) described augite exsolution lamellae along 001 within pigeonite and vice versa and deduced that the Shergotty pyroxenes had cooled at 0.002°C/h at temperatures of 1100-800°C: a figure comparable with large terrestrial intrusives and consistent with the relatively slow cooling at high temperatures suggested by the presence of symplectites in basaltic shergottites. We discuss the depth of origin of the shergottites in relation to magma water content in a later section.

McSween et al. (1996) showed that the zonation patterns of Fe-enrichment towards the margins of pyroxene grains were similar to those of lunar pyroxenes that had crystallised from melts. Kring et al. 2003 also showed that QUE94201 pyroxene core compositions were consistent with formation from a melt with the composition of the bulk sample. Thus QUE94201 is regarded as a bulk melt composition (e.g. McSween et al. 1996; Wadhwa et al. 1998) whereas most of the other shergottites contain some cumulus pyroxene (Hale et al. 1999). Apart from QUE94201 and Los Angeles the shergottites also contain too high a proportion of pyroxene to have formed as 100% melts (McSween, 1994). We discuss the phase relations of the SNCs and their associated melts in later sections. Evidence that Mg-rich pyroxene cores in most of the basaltic shergottites accumulated rather than crystallised in-situ has often been taken from the preferred orientations of the pyroxene grains which define a layered fabric. However, although the basaltic shergottites are often described as cumulates there is no compositional evidence that crystal-melt segregation or accumulation of phenocrysts had any major effect on the rock compositions. Hale et al. (1999) calculated the proportion of homogenous Mg-rich cores of pyroxene within Zagami and Shergotty (14 and 19%) and suggested that this was the proportion of cumulus grains.

Mittlefehldt (1999) suggested that EET79001 lithology A was an impact melt with a hybrid composition derived from basaltic lithology B and lherzolitic cumulates. However, the impact melt model for EET79001B requires a remarkable coincidence between the age of the impact, i.e. the crystallization age of lithology B, 174±3 Ma (Nyquist et al. 2001b), and the 175 (±~10) Ma mode among regular igneous ages for shergottites. Warren and Kallemeyn (1997) cited additional weaknesses of the impact melt hypothesis, including siderophile (Au) data, the limited variety of clast types within the putative impact melt, and the gradational nature of the “A” to “B” contact; a constraint reinforced by the detailed petrologic study of van Niekerk et al. (2004).
Olivine-rich shergottites

Eight of the basaltic shergottites contain up to 7-29% large (1-3 mm) olivine grains (Table 1, Fig. 1) and have been described as olivine-phyric shergottites or picritic shergottites (Gnos et al. 2002, Goodrich, 2003). This includes lithology A of EETA79001. These shergottites also have a higher bulk Mg# range of 59-68 (Goodrich, 2003) than the basaltic shergottites and contain chromite, rather than Ti-magnetite and ilmenite, as the main oxide phases (Table 3). The olivine in SaU005 is Fo02-72 (this study), in EETA79001A it is Fo33-76 (Goodrich, 2003); the larger grains having the more Mg-rich compositions. The most Mg-rich olivine of the olivine-rich shergottites is found within Y-980459 (Fo84), (Greshake et al. 2004). One of the characteristics of the SNCs is that they have relatively oxidised mineral assemblages compared to other meteorite groups. For instance chromite cores in SaU005, DaG476 and EET79001A have 100Cr/(Cr+Al) atomic ratios of 74-87, 100Mg/(Mg+Fe 2+) 12-36 and 100Fe 3+/(Fe3++Cr+Al) 0-5 (Bridges and Grady, 2001). Pyroxenes in the olivine-phyric shergottites are more Mg-rich than in the basaltic shergottites: SaU005 and DaG476 are En66-77 Wo7-33 Fs16-31 (Fig. 2).

McSween and Jarosewich (1993) and McSween (1994) proposed that the olivine grains were xenocrysts resulting from the mixing of peridotitic material and basaltic shergottites. However, in a study of their textures, Goodrich (2003) noted that many of the large olivine grains were compositionally zoned and subhedral, describing them as predominantly phenocrysts, which may however have been brought in to the magma rather than crystallising in-situ. Uniquely for the shergottites, Y-980459 contains no plagioclase or maskelynite and was inferred to have cooled rapidly upon extrusion onto the martian surface after crystallisation of Mg-rich cumulus phases in an underlying magma chamber (Greshake et al. 2004).

Peridotitic (lherzolite) Shergottites

The 6 peridotitic shergottites have similar mineral assemblages to the olivine-phyric basaltic shergottites but contain <10 vol. % maskelynite, 40-60% olivine, 9-25% pigeonite and augite; with bulk Mg# ~70 (e.g. Goodrich 2003; McSween et al. 1979, Mason, 1981). They are cumulates with euhedral olivines typically poikilitically enclosed by pigeonites or augites up to 5 mm across. The pigeonites sometimes contain exsolutions of orthopyroxene and high-Ca pyroxene. Some peridotitic shergottites also show a clearly defined alignment of cumulate grains (Berkley and Keil, 1981).
Mineral compositions of the peridotites overlap those of olivine-phric shergottites, with a range of olivine compositions of Fo$_{60-76}$ (McSween and Treiman, 1998). The peridotitic shergottites are often referred to as the lherzolitic shergottites but as they lack significant quantities of orthopyroxene the term peridotite is more accurate. Chromite has similar compositional ranges and Fe$^{3+}$ contents to those in the olivine-phric shergottites.

**Nakhlites and Chassigny**

The 7 nakhlites are cumulate olivine-clinopyroxenites composed of augite, Fe-rich olivine and mesostasis (Tables 1, 3, Fig. 3). One of the nakhlites MIL03346 is an olivine-free clinopyroxenite. Lentz et al. (1999) published modal abundances of Nakhla, Lafayette and Governador Valadares showing that they contained similar proportions of the main mineral phases: olivine 5-20%, pyroxene 69-85%, mesostasis 5-13%. An average olivine content based on our own studies of nakhlite sections (Nakhla, Lafayette, Governador Valadares, Y000593) is consistently ~10%. However, NWA817 has a distinctly higher proportion of mesostasis than the other nakhlites (20%, Sautter et al. 2002; Mikouchi et al. 2003). Mesostasis in the nakhlites is composed of plagioclase and alkali feldspar dendritic grains, pyroxene dendrites, glassy mesostasis, Ti-magnetite and ilmenite, silica polymorphs, chlorapatite and sulphides together with occasional baddeleyite and secondary mineral assemblages e.g. siderite (Bridges and Grady, 2000). The presence of silica polymorphs shows the silica-oversaturated nature of the mesostasis. Unlike Chassigny (see below) the augite cores (En$_{0.37-0.62}$Wo$_{0.37-0.43}$Fs$_{0.24-0.41}$) and olivine (Fo$_{14-44}$) studied here have not been equilibrated to a great degree (Fig. 4). The augite rims are zoned to Fe-rich compositions, hedenbergite in NWA817. NWA817 olivine also shows the greatest range of olivine compositions of the nakhlite samples, including some zonation within individual grains, from Fo$_{14-44}$ (Sautter et al. 2002).

The presence of radiating plagioclase dendrites in the mesostasis (Fig. 3) may require that the final cooling of interstitial liquid took place rapidly in a quenching event, subsequent to the slower cooling associated with accumulation of the augite and olivine grains. Magmatic or melt inclusions (predominantly fine-grained pyroxene and feldspar, glass, apatite, ilmenite intergrowths) in the nakhlites, especially within olivines, have been used to reconstruct parent melt compositions e.g. Treiman (1993) and we consider this further in relation to phase equilibria of martian basalts in a later section.
Chassigny is a cumulate dunite, having >90% olivine. The remainder is composed of augite and low-Ca pyroxene (occasionally poikilitically enclosing the olivine), plagioclase, minor orthoclase and chromite and melt inclusions with H$_2$O-bearing kaersutite amphibole, biotite and glass (Floran et al. 1978, Johnson et al. 1991). The olivine (Fo$_{68-71}$) and chromite (Mg$\# = 12-19$) have relatively narrow compositional ranges showing the equilibration at 1150 – 1230°C (Wadhwa & Crozaz, 1995). However there is evidence (Floran et al. 1978) for compositional heterogeneity in the chromite analyses. This sample has similar crystallization and ejection ages to the nakhlites and so is often regarded as having originated as part of the same set of igneous rocks on Mars. Both the nakhlites and Chassigny did form from LREE-enriched magmas but Wadhwa and Crozaz (1995) suggested that the differing slopes of the REE profiles are not consistent with formation from the same melt. Despite the presumed close proximity of their origin (on the basis of their very close ejection ages) there is no clear model to relate the magmatic history of the nakhlites and Chassigny.

**ALH84001**

ALH84001 is an orthopyroxenite with a cataclastic texture. Orthopyroxene grains are up to 6 mm in diameter occupying 97% of the rock with maskelynite (mainly An$_{31-37}$), chromite, augite, apatite and pyrite representing the remnants of a small amount of trapped interstitial melt. Mittlefehldt (1994) proposed that it was a cumulate on the basis of its near monomineralic, equilibrated nature, coarse grain size and homogenous pyroxene composition (mean is Fs$_{28}$Wo$_3$En$_{69}$). Its small degree of trapped melt means that it can be considered an adcumulate. Chromite in ALH84001 has 100Fe$^{3+}/$(Fe$^{3+}$+Cr+Al) = 4.5-10 (Bridges and Grady, 2001). ALH84001 is particularly famous for its secondary carbonate ‘rosettes’ e.g. Mittlefehldt (1994), McKay et al. (1996); Bridges et al. (2001).

**Physical Parameters**

The calculated room-temperature densities for SNC meteorites vary from 3.59 g/cm$^3$ for the Chassigny dunite to 3.18 g/cm$^3$ for EET79001B basaltic shergottite (Lodders, 1998). Peridotite LEW88516 is 3.39 g/cm$^3$ and Nakhla is 3.33 g/cm$^3$. The high FeO contents of the SNC melts are associated with high magma densities relative to a martian crustal density of ~3.0 g/cm$^3$ – allowing the ponding of large magma bodies. This and low viscosities are both consistent with the scale and morphology of many observed martian lava flows. In Table 4 we show our calculated melt densities at a liquidus temperature of 1200°C and 1.5 kbar for the basaltic shergottites, and the *Pathfinder, Spirit*
and Opportunity results. The low-pressure density of a silicate melt can be calculated to good precision based on its composition and the partial molar volumes of its constituent oxides (Lange and Carmichael, 1990). The 1.5 kbar pressure is equivalent to 10 km depth (Turcotte and Schubert, 1982) i.e. upper to mid crustal thickness for much of Mars. These data show the expected lower densities for the basaltic-andesitic-like rocks from Pathfinder e.g. 2.61-2.88 g cm\(^{-3}\) compared to the basalts: Humphrey’s melt is 2.85 g cm\(^{-3}\) and the basaltic shergottite predicted melts are 2.82-2.90 g cm\(^{-3}\). The nakhlite parent melt NK93 (Treiman, 1993) has a density of 2.80 g cm\(^{-3}\). Further details of the melt density calculations are available from the authors. The estimated viscosity range for crystal-free, hydrous SNC melts is 0.1-30 N sm\(^{-2}\) (Harvey et al. 1993). However the presence of entrained cumulate crystals – as are seen in the shergottites - may act to increase the viscosity by a factor of 10 (Basaltic Volcanism Study Project, 1981).

**The SNCs: Geochemistry**

Geochemical classification of the shergottites

As discussed above, shergottites are usually subclassified on mineralogical grounds. A separate yet equally valid way of subclassifying shergottites is based on REE geochemistry (Fig. 5, Table 5). There are again three groups, which we here define as highly depleted (HD) shergottites, moderately depleted (MD) shergottites, and slightly depleted (SD) shergottites. Even the SD shergottites (which constitute the most numerous subclass) are in some respects depleted vs. chondrites, but the depletions are only very mild in comparison to HD shergottites. These geochemical subclasses correlate only loosely with the mineralogical subclasses. Most of the HD shergottites are olivine-phyric, but QUE94201 is a basaltic shergottite (Wadhwa et al. 1998; Kring et al. 2003). Most of the MD shergottites are cumulate peridotites; but EET79001 contains both basaltic and olivine-phyric MD lithologies. Most of the SD shergottites are basaltic without conspicuous phenocrysts; except NWA1068/1110 which is olivine-phyric (Barrat et al. 2002).

“Depleted” refers to the many elements that behave incompatibly with respect to major minerals during igneous differentiation. Exemplary incompatible elements include Th, U, Ba, and La. The ratio La/Lu is particularly diagnostic. Fig. 5 shows CI chondrite-normalized REE patterns for a representative selection of shergottites. The depleted subtype have La/Lu consistently close to 0.12 times CI, whereas the SD subtype have La/Lu consistently close to 1.0 times CI. Despite their cumulative...
nature, the peridotitic shergottites have REE patterns (including La/Lu) lower but roughly parallel to the patterns of their parent melts (Wadhwa et al. 1994). For comparison, mid-ocean-ridge basalts, which represent the major depleted reservoir within the Earth’s mantle, typically have La/Lu ~0.6 × CI (Basaltic Volcanism Study Project, 1981). In a later section we demonstrate that these trace-element signatures of highly diverse extents of depletion are paralleled by variations in initial $\text{Sr}^{87}/\text{Sr}^{86}$ ratios (Borg et al. 2002). The systematic depletion variations among shergottites tend to correlate, but only in a loose way, with key mineralogical traits. To assume that depletion/enrichment is a reliable predictor of mineralogical subclass or vice versa (Herd, 2003; Goodrich, 2003) can be a mistake.

**Whole Rock and Melt Compositions**

The compositions of the SNC group are similar to terrestrial igneous rocks in some respects. Stolper (1979) demonstrated that the siderophile and volatile trace element abundances of basaltic shergottites differ from those of terrestrial basalts by less than a factor of ten. The clearest differences in major element composition between the SNC meteorites and terrestrial igneous rocks are the high Fe contents of the martian samples. Fe, expressed as FeO, ranges from 17.3 wt% for ALH84001 to 27 wt% for Chassigny. QUE94201 has 18.5 wt% FeO (Table 4). In contrast typical terrestrial ocean floor basalts and andesites have <10 wt% FeO. The SNCs have also often been considered to show Al-depletion, having Al$_2$O$_3$ contents of 0.7 wt% (Chassigny dunite) to 11 wt% for the basaltic shergottites QUE94201 and EETA79001B (Lodders, 1998). In Fig. 6 the plots of CaO v. MgO and Al$_2$O$_3$ v. MgO show the generally low Al$_2$O$_3$ contents and similar CaO of most the SNCs compared to terrestrial basalts but the *Pathfinder* basaltic-andesites (discussed below) and *Spirit, Opportunity* basalt analyses do not exhibit notable Al-depletion. In Table 4, the *Pathfinder, Opportunity* and *Spirit* results show Al$_2$O$_3$/CaO ratios similar to terrestrial basalts.

The bulk contents of H$_2$O in the SNC meteorites are low e.g. 300 µg/g for Shergotty (Lodders, 1998). The nakhlites have higher H$_2$O contents but these are a result of alteration processes on the martian surface. Dann et al. (2001) reported the results of phase equilibria experiments on Shergotty compositions. They suggested that elevated H$_2$O pressures ~1.8 wt% were necessary for their melts to crystallise pigeonite and augite as liquidus phases at magmatic temperatures 1000 – 1200°C. If this is the case then the shergottite magma must have lost the majority of its water during ascent and prior to final crystallization. This melt water content is anomalous because the H$_2$O content of the martian mantle is considered to be <300 µg/g (Lodders & Fegley, 1997) and therefore some form of crustal...
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contamination associated with the addition of water into the melt has been considered necessary by various authors. Crystallisation of the basaltic shergottite pyroxene cores commenced slowly at depth from H₂O-bearing melts (e.g. 4 - 6 km, 1.8 wt% H₂O) and was followed by more rapid crystallisation of the Fe-rich rims – and in the case of Y980459 the entire matrix – in a near-surface intrusive or extrusive setting (Greshake and Stoffler, 2004; Lentz et al. 2001; Beck et al. 2004).

The SNC group contains both melt compositions e.g. QUE94201 and cumulates e.g. the nakhlites, Shergotty. In the following sections we consider melt major element compositions and relate them to the Pathfinder, Spirit and Opportunity in-situ analyses of igneous rocks.

**Basalts and Basaltic Andesites on Mars:** The recalibration and processing of the Pathfinder data to remove the effects of dust cover are described in Wänke et al. (2001) and Foley et al. (2003). Foley et al. used the combined alpha, proton and X-ray modes and so this may be considered the more comprehensive data set. They demonstrated that the Pathfinder rock analyses contained up to 4 wt% H₂O, suggesting that alteration rinds were sampled. The Pathfinder soil-free rock recalculated analysis of Foley et al. has high SiO₂ 57.7 wt%, high K₂O 1.20 wt% and low MgO 0.8 wt% corresponding to terrestrial andesites or basaltic andesites (Figs. 6, 7). However, the direct evidence for olivine and pyroxene grains (i.e. igneous mineralogy) identified in basalts at Gusev by the Spirit Lander, is lacking for the Pathfinder analyses. Therefore the classification of Pathfinder rocks as basaltic andesites still has some uncertainty, with the possibility remaining that the high alkali contents are a result of unidentified contamination by dust or alteration material or even that the rocks are not igneous (Foley et al. 2003; McSween et al. 1999).

In contrast the basaltic shergottites (Lodders, 1998) are ~48-51 wt% SiO₂, <0.2 wt% K₂O and 6-16 wt% MgO, and, despite their low Al₂O₃ and high FeO contents discussed previously, have recognizably basaltic compositions. Mixing of 28% of the shergottite component with 54% of the Pathfinder, andesitic component together with lesser amounts of Mg-sulphates and ferric oxides fits the composition of martian soil analysed by the Viking and Pathfinder landers (Wänke et al. 2001).

Primitive and shergottite-like basalts have now also been identified at the MER 2004 landing sites (McSween & Jolliff, 2004; McSween et al. 2004; Zipfel et al. 2004). ‘Bounce Rock’ at Meridiani Planum has 51 wt% SiO₂, 0.1 wt% K₂O, 6.5 wt% MgO (Table 4, Figs. 6, 7) and is thus similar to the basaltic shergottites. However, analyses of Gusev rocks – some of which contain large olivine crystals – have lower CaO (7.5-7.7 wt%) and higher Na₂O + K₂O contents than the basaltic shergottites and a
more picritic composition. Mössbauer and Mini-TES spectra confirm the presence of olivine, magnetite, and probably pyroxene in the Spirit-Gusev analyses (McSween et al. 2004).

The high MgO, and low CaO, TiO₂, Al₂O₃ contents of the peridotites (Fig. 6a-c) reflect their high modal abundances of olivine grains. Similarly the lower MgO contents of the basaltic shergottites reflect the absence of significant proportions of olivine. The olivine-phyric shergottites are intermediate in composition. The simplest explanation for the trends of negative correlation between MgO and CaO, TiO₂, Al₂O₃ for the meteorite analyses is that the olivine-phyric shergottites are a mixture of peridotitic xenocrysts with shergottitic melt. This is the model advocated by McSween (1994) and McSween and Treiman (1998). However, as we demonstrate using La/Lu and initial ⁸⁷Sr/⁸⁶Sr ratios of the shergottites in a later section, this is inconsistent with the geochemistry of the peridotitic shergottites (all MD) and olivine-phyric shergottites (consistently HD). Therefore even if the olivine grains within the olivine-phyric shergottites were brought in from peridotitic sources, in order to explain the possible mixing, they were not the same suite of peridotites sampled within the SNC meteorites.

Goodrich (2003) proposed a more complex model than that of McSween and Treiman (1998). She noted that the parent magma for the large olivine grains in SaU005 (predicted from melt inclusions in chromite) was too CaO-rich to be the same as that associated with the peridotites. Therefore she proposed that in addition to some material (the most Mg-rich xenocrysts or phenocrysts) that was mixed in from unidentified peridotitic sources, most of the large olivine, low-Ca pyroxene and chromite grains were derived from a separate, olivine-saturated basaltic melt. She further speculated that such a melt could have been parental to the basaltic shergottites. This idea is consistent with the work of Stolper and McSween (1979) who demonstrated with experimental phase equilibria studies of the basaltic shergottites that the primary melts from which the shergottitic melt evolved was olivine-saturated.

Fig. 8 shows the phase relations of the SNCs and their associated melts in si-ol-plag space projected from wollastonite using the oxygen unit plotting parameters (equivalent to volume) of Longhi and Pan, (1988); Longhi, (1991). The cumulate nakhlites, Chassigny and peridotite shergottites plot towards the olivine apex. Some of the samples e.g. Bounce Rock and Al-rich SD shergottites plot close to the mafic-plag cotectics, consistent with a basaltic melt rather than primarily cumulate origin. The calculated nakhlite melt of Treiman (1993) lies closest to the low-Ca px – ol cotectic consistent with
the late crystallisation of plagioclase in the mesostasis of the nakhlites. The low plagioclase contents of the SNCs compared to the Pathfinder analyses is evident.

There is no clear link between the basaltic andesite compositions from Pathfinder, and the basalts from shergottites and Spirit and Opportunity. However, Dann et al. (2001) showed that crystallization of shergottitic melts under water-saturated conditions can lead to FeO and Al₂O₃ depletion and SiO₂ enrichment to produce andesitic-like melts. In a similar set of experiments, Minitti and Rutherford (2000) showed that 1-2 wt% H₂O in the experimental charges was necessary to produce enough melt (40 vol.%) of andesitic composition to separate from the crystals.

Geochemical evidence for the importance of water in the evolution of shergottites is given by Li geochemistry. Lentz et al. (2001) and Beck et al. (2004) showed that some pyroxenes in shergottites exhibited a decrease in Li abundances from cores to rims: this was attributed to exsolution of water from the magma at depth (4-6 km). The more Li-enriched rims of some pyroxenes were attributed to their crystallisation near the martian surface (<0.5 km).

The terms basaltic andesites and andesites have become widely used in the context of martian geology, and so we continue to use the terms in this paper. However we note that for terrestrial rocks the name ‘icelandite’ is generally applied to those rocks of andesitic composition which formed in a nonorogenic setting. The (Na₂O + K₂O) and SiO₂ contents of icelandites and andesites are identical (Middlemost, 1985) and as no orogenic processes have ever been convincingly described on Mars the term andesite may be misleading for a martian rock (McSween et al. 1999).

Nakhlite and Chassigny Cumulates and Parental Melts

The parental melt compositions of the nakhlites have been studied in a variety of ways. Treiman (1993) used electron microprobe analyses of melt inclusions and deduced that the parental melt was basaltic, rich in Fe, and poor in Al compared to terrestrial basalts (Table 4). However, Lenz et al. (1999) suggested that the melt had textural and compositional affinities to an Archean ultramafic lava flow (Theo’s Flow) with the exception of lower Al₂O₃ contents and Al₂O₃/CaO ratios which reflected the Al-depleted martian mantle (e.g. Longhi, 2002). Theo’s flow formed within a 120 m thick lava flow, with settling of cumulus pyroxene grains and continued growth in a growing cumulate pile and this may be analogous to the formation conditions of the nakhlites. Melt inclusions within Chassigny olivine have also been used to calculate parental melt compositions. Like the nakhlites the calculated
melt is basaltic but FeO-rich and Al$_2$O$_3$-poor with an inferred 1.5 wt% H$_2$O dissolved in the melt prior to kaersutite crystallization (Johnson et al. 1991).

In this section we show that the nakhlites formed as cumulates followed by the trapping of LREE > HREE basaltic (albeit an unusually mafic and low-Al composition) melt. As shown previously, there is an almost complete overlap in augite compositions between the nakhlites (En$_{0.37-0.62}$Wo$_{0.37-0.43}$Fs$_{0.24-0.41}$) but NWA817 pyroxene extends to significantly more Fe-rich compositions (Wo$_{22}$En$_{22.38}$Fs$_{50.36}$, Sautter et al. 2002, Fig. 4) suggesting that this nakhlite did crystallise from a more fractionated melt.

The rims (e.g. ≤20 µm in Governador Valadares, Fig. 9) are also more Fe-rich and Mg-poor than the cores. For instance, Harvey and McSween (1992) analysed a series of core-rim augite pairs in Nakhla and showed that the rims were up to Fs$_{0.6}$ in one analysis and mainly around Fs$_{0.4}$. The olivine grains within the nakhlites, apart from NWA817, are Fo$_{29-38}$. NWA817 olivine is Fo$_{15-42}$ (Mikouchi et al. 2003) and shows zonation from core to Fe-rich rims. Thus the nakhlite olivine and the augite are not entirely in Mg-Fe equilibrium. This is shown in Fig. 9a, b, X-ray maps (Fe K$\alpha$ and Mg K$\alpha$) of Governador Valadares. An explanation for this feature was provided by Harvey and McSween (1992), who suggested that olivine equilibrated more rapidly than pyroxene. However, as for instance Nakhla has a bulk Mg# of 51 (Table 1, Fig. 4) the olivine has clearly not equilibrated to the bulk composition of the rock. An alternative explanation of the disequilibrium may be given by the microscopic textures. Olivine has crystallised after and around the augite cores but before their Fe-rich rims. Olivine compositions are nearer to equilibrium with the final melt, rather than with the first-formed pyroxene.

The late crystallization of the olivine will also have acted to drive the composition of remaining trapped melt towards silica-oversaturated compositions. The mesostasis (which contains silica polymorphs) and the Fe-rich augite rims are the crystallised remnants of trapped melt. The proportions of trapped melt in the nakhlites are thus ≤13%, except for NWA817 which may be higher (Mikouchi et al. 2003). The low proportion of trapped melt is consistent with an origin for the augite grains primarily through cumulus growth, maintaining the composition of the augite, until cumulus growth ends and the trapped melt and Fe-rich rims crystallised. The mineral zonation preserved in NWA817 suggest that it underwent the least thermal equilibration of the nakhlites as a result of rapid cooling, at the top of the pile of nakhlite cumulates.

REE and fractional crystallisation in the nakhlites: The petrographic model described in the previous section can be quantified through use of trace element compositions of the nakhlites. Nakhla
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(BM1913, 26) and Y000593 have nearly identical relative abundances of the REE e.g. La/Lu = 3.5, 4.0 (Table 6, Fig. 10) Nakhla has 4.5 – 1.3 x CI REE with Y000593 having slightly higher measured abundances 6.9 – 1.7 x CI. The range of trace element abundances of nakhlite augite and mesostasis are shown in Fig. 11. The cumulus augite cores for the 5 nakhlites analysed have indistinguishable REE abundances with 0.5-5 x CI and La/Lu ~0.75, without discernible Eu anomalies. Mesostasis analyses – containing a fine-grained mixture of glass, silica, feldspar, phosphate – reflect trapped melt compositions. They are LREE-enriched with ≤400 x CI abundances and La/Lu ~ 14 - 30, generally showing smooth patterns. The lower abundances e.g. 100 x CI La – 7 x CI Lu for Y000593 are closest to the initial melt composition. This potential parental melt composition is similar to that calculated for Nakhla by Wadhwa & Crozaz (1995) and Wadhwa & Crozaz (2003) on the basis of early-formed augite core REE abundances and a (basaltic) melt inclusion within NWA998. They suggested that the Nakhla parental melt was <60 x CI for La with La/Lu ~10. The LREE > HREE smooth abundance patterns for the predicted parental melt are similar for the mesostasis with the lowest REE abundances and melt inclusion data of Wadhwa and Crozaz (2003). Here we use the Wadwha and Crozaz data in Fig. 11 to model the formation of the augite-olivine cumulates because it has the lowest REE abundances, and so is consistent with an early, parental melt.

We use the Rayleigh fractionation equation [1], where C_L and C_o are REE concentrations within the evolving melt and original parental melt. We use a value of F (fraction of melt remaining) of 0.6 because this value is typical for ultramafic magmatic crystallization products (e.g. Snyder et al. 1995). The solid-melt trace element distribution coefficient D [2] was calculated as the weighted mean for an unfractionated olivine (10%)-augite (90%) assemblage. This is taken to be the cumulate pile prior to final crystallization and trapping of mesostasis. Appropriate values of D for olivine and augite were taken from Papike et al. (1996).

\[
\frac{C_L}{C_o} = F^{(D-1)}
\]

\[
D = \bar{a} w_i K_{pi}
\]

The REE compositions formed from fractional crystallization of the Wadwha and Crozaz (2003) parent melt are plotted in Fig. 10. It is 100 – 12 x CI with La/Lu ~ 9 and a smooth abundance pattern. Addition of between 5 and 20% of this melt to the ol-augite cumulate is consistent with the trapping of
melt within the mesostasis. The profiles so formed have similar REE abundances and LREE/HREE fractionation to the nakhlites’ whole samples. The figure of 10% is similar to the proportions of mesostasis in Nakhla and Y000593 and 20% is similar to the proportion of mesostasis in NWA817 (Mikouchi et al. 2003) suggesting that our model can account for the REE abundances in the nakhlites. Fig. 12 (after Mikouchi et al. 2003) summarises the formation of the nakhlite cumulates.

**Planetary Differentiation and Origin of the SNC melts**

The evidence for Fe-rich SNCs, the *Viking* and *Pathfinder* analyses and basalts observed on the surface with low viscosity have led to the widespread assumption (e.g. McSween, 1994) that the martian mantle is Fe-rich (Table 4). The mass of the core has been estimated at 22% of the total Mars mass by Dreibus and Wänke (1985) based on SNC data. The Mars core is also assumed to be S-rich because of the depletion of chalcophile elements in SNCs (e.g. Dreibus and Wänke, 1985). This small core (e.g. the Earth’s is 30% by mass) is associated with a correspondingly high FeO content in the Mars mantle: 19 wt% (Mg# ~0.75) compared to 8 wt% for the Earth (Table 4). Treiman et al. (1987), considering the calculated abundances of siderophile and chalcophile elements within the mantle SNC source region, suggested a higher range from 25-35% of the mass of Mars was within the core. However the moment of inertia for Mars is now well constrained through tracking of the *Pathfinder* lander (Folkner et al. 1997) at 0.366. Bertka and Fei (1998) demonstrated that such a figure was consistent with the Fe contents suggested by Dreibus and Wänke (1985) and Lodders and Fegley (1997).

**Evolution of the martian interior: late accretion and core formation**

Important insight into the evolution of the martian deep interior can be obtained through highly siderophile elements (HSE), so-called because they partition strongly into metal in preference to silicate (e.g. Borisov et al. 1995). This behaviour makes HSE sensitive recorders of core formation and of metal-silicate interactions in general. Earth’s upper mantle contains moderately high levels of HSE (~0.008 × CI chondrites) that are often interpreted as evidence for “tardy” accretion of ~0.5 wt% of the Earth as a chondritic “late veneer,” subsequent to the primordial brief (or non-existent?) period of efficient core-mantle equilibration (e.g. Schmidt et al. 2003; Walker et al. 2004). The time-integrated history of accretion is one of the great unknowns of planetology, but data for lunar samples (recent
review: Warren, 2004) and achondritic meteorites (Bogard, 1995) indicate there is a significant “tail”
effect, with a final spurt intense enough to be popularly known as a “terminal cataclysm” at ~3.85 Ga.
An alternative interpretation (e.g. Righter and Drake, 1997) is that during Earth’s core formation
extremely high pressures greatly moderated the HSE metal/silicate partition coefficients.

Of the many elements that have frequently been determined in martian meteorites, the most
thoroughly siderophile are Os and Ir. In pure mafic-silicate systems, these elements show compatible
behaviour, such that they tend to correlate (albeit not very tightly or linearly) with MgO (e.g. Naldrett
and Barnes, 1986). A plot of MgO vs. Ir (Fig. 13) shows the correlation among martian meteorites
(using Os in lieu of Ir would result in a nearly identical distribution). The ancient cumulate
orthopyroxenite falls well away from the general trend but the remaining samples show a distribution
remarkably similar to that of mafic-igneous rocks from Earth. Both the Mars and Earth trends are
greatly Ir-enriched in comparison with the overall trend of the lunar data. In evaluating the relative
positions of the trends on Fig. 13, it should be recalled that Mg# is probably roughly 2× lower in the
martian mantle than in the mantles of the other two bodies; and this Mg# disparity is probably
associated with slightly lower MgO, for an otherwise comparable martian rock, in comparison to a
terrestrial or lunar counterpart. However, this is a detail, given the factor of 5 range in MgO among the
martian meteorites. The crystalline lunar mare basalts are not obviously Ir-poor in comparison with
shergottitic basalts, but the MgO-rich nonmare cumulates 67667 and 72417 indicate a major downward
displacement of the lunar trend, and recent careful analyses (isotopic as well as elemental) of two types
of mare-pyroclastic glass by Walker et al. (2004) indicate a similar, if not larger, relative depletion of
HSE in the lunar mantle, vs. the mantles of Mars and Earth.

These trends indicate that just like Earth, Mars retained levels of HSE in its mantle equivalent to
slightly less than 1 wt% of CI-chondritic component. In contrast, the much smaller Moon managed to
achieve and sustain a relatively efficient depletion of HSE from its mantle. The similarity between the
HSE depletions of the martian and terrestrial mantles is a remarkable coincidence. It has occasionally
been claimed (or implied) that Earth’s mantle siderophile “discrepancy” is “unique to the Earth among
all documented planetary objects” (Jana and Walker, 1997); and that some peculiar trait of large
planets, such as extreme pressure (e.g. Righter and Drake, 1997) might account for this “unique”
evolution. It now seems clear (cf. Warren et al. 1999) that HSE behaviour in the core-mantle evolution
of Mars closely paralleled HSE behaviour during evolution of Earth. Yet the pressure at the
mantle/core boundary is very different in Mars (~23 GPa) versus Earth (~136 GPa). Rather than an effect of pressure on HSE partitioning behaviour, the trends in Fig. 13 might be functions of the efficiency with which late-accreted chondritic materials were stirred down into the mantle (“late” here means after the interior has cooled, from a primordial largely molten state, to the point where the mantle, being mostly solid, no longer effectively equilibrates with the core). Even after reaching a largely solid state, the Earth’s mantle (as is well known) continued to convect in a style that, over time, mixes surface material with the deep interior. The post-magma-ocean Moon is and probably always has been incapable of this type of down-stirring. There are reasons to doubt that Mars ever sustained a tectonic style with Earth-like major involvement of surface plates in mantle convection (e.g. Warren, 1993). But the similarity between the Mars and Earth trends in Fig. 13 suggests that at least briefly in its post-core/mantle-equilibration evolution, Mars sustained a style of convection/tectonism vigorous enough for it to manage (with an efficiency closer to 100% than to 0%) a sweeping of near-surface “late” accretion matter down into its mantle.

**Magma Sources**

The diversity of depletion characteristics among shergottites (Fig. 5, 14) shown in our geochemical classification has often been interpreted as a manifestation of varying extents of assimilative mixing between primary, HD-type basalt and a putative enriched (high La/Lu, REE-rich, Sr-rich, etc.) material (e.g. Nyquist et al. 2001a, b; Borg et al. 2002). The trace-element signatures of highly diverse extents of depletion do parallel isotopic variations. For example, initial \(^{87}\text{Sr}^{86}\text{Sr}\) ratio, normalized to the typical shergottite age of 175 Ma, is consistently close to 0.701 among highly depleted shergottites, close to 0.722 among SD shergottites, and close to 0.711 among the MD shergottites (Fig. 14). A very similar pattern is manifested using \(\varepsilon_{\text{Nd}}\) (at 175 Ma) in lieu of initial \(^{87}\text{Sr}^{86}\text{Sr}\) (Borg et al. 2002).

The setting for the assimilative mixing and the detailed composition of the enriched component are speculative; the assimilated material is often referred to simply as “crust.” Nyquist et al. (2001b) have proposed a roughly quantitative model in terms of Sr and Nd, including isotopic ratios. However, these authors did not model La/Lu (or any analogous ratio such as La/Yb). The red curve in Fig. 14 shows a mixing model based on the assumptions of Nyquist et al. (2001b) for Nd, Sr and \(^{87}\text{Sr}^{86}\text{Sr}\) in the crust and HD-basalt components, augmented with an assumption that the La/Lu of the crust component was equivalent to the lunar (KREEP) crust in terms of REE pattern, i.e. that it had La/Lu = 2.28 × CI.
chondrites (Warren, 2004). This extension of the Nyquist et al. (2001b) model yields a marginally satisfactory fit to the array of shergottite compositions.

It should be noted, however, that the Nyquist et al. (2001b) model yielded (by extrapolation from all the other assumed parameters) a dubiously low Sr concentration for the HD-basalt component (specifically, 11.5-26.2 µg/g; note: only the average of 19 µg has been employed for our Fig. 14), which translates into relatively high (crust-dominated) $^{87}\text{Sr}/^{86}\text{Sr}$ at any given La/Lu position along the Fig. 14 mixing curve. No known shergottite analysis has Sr <20 µg/g. The mass-weighted mean of two precise analyses for QUE94201 (Borg et al. 1997) is 48 µg/g, and the only other applicable result (most HD shergottites have suffered warm-desert weathering) is 20 µg/g for Y980459 (Shih et al. 2003. Fig. 14 also includes a set of three curves based on the more realistic assumption that the average HD composition has Sr = 35 µg/g. The thick black curve is otherwise analogous to the red (19 µg/g) one, but the other two 35 µg/g curves illustrate the sensitivity of the mixing parabolas to the assumed La/Nd and Lu/Nd (and thus La/Lu) ratios of the enriched component. In the case of the upper curve, the enriched component’s REE pattern is assumed to parallel the pattern of the Earth’s continental crust (Taylor, 1992), with La/Lu = 6.6 × CI. In the case of the lower curve, the enriched component’s REE are assumed to be unfractionated (“flat”) with respect to La/Lu. In all these 35 µg/g models, the proportion of the enriched component that has to be added to the HD end-member to reach the SD shergottites is ~ 17 wt%; in the 19 µg/g model, it is (perhaps more realistically, from a thermo-physical standpoint) ~10 wt%. Note that only the model with flat La/Lu comes close to matching the combination of low La/Lu and relatively high initial $^{87}\text{Sr}/^{86}\text{Sr}$ observed in the two distinct (but similar) EET79001 lithologies. An approximately flat REE pattern for an otherwise highly enriched component, while conceivable, is hardly expected as a typical outcome from planetary igneous differentiation.

Another parameter that might be at play is the enriched component’s Sr level, but the 204 µg/g assumed by Nyquist et al. agrees with an estimate of 180 µg/g, based on martian meteorite Sr/K systematics coupled with the Pathfinder soil K concentration, by Dreibus and Jagoutz (2003), albeit new data from the Spirit and Opportunity sites show only half as much soil K as was found by Pathfinder (Rieder et al. 2004). Appeal might also be made to a higher $^{87}\text{Sr}/^{86}\text{Sr}$ in the crustal component. However, Dreibus and Jagoutz (2003) argue that the $^{87}\text{Sr}/^{86}\text{Sr}$ assumed by Nyquist et al. (2001b) was already too high to represent the average crust.
Borg et al. (2002) extended the model of Nyquist et al. (2001b) by proposing that the crustal component was assimilated in an episode of assimilation during fractional crystallization (AFC). However, this variant of the assimilation model seems even less well-suited to account for the La/Lu constraints (Fig. 14), since the proposed fractional crystallization (including major high-Ca pyroxene) would enhance La/Lu while having no (direct) effect on the $^{87}\text{Sr}/^{86}\text{Sr}$ ratio.

In summary, it appears that a 2-component mixing model (Nyquist et al. 2001b; Borg et al. 2002) yields a marginal fit to the MD shergottite data, but only if La/Lu in the enriched component is assumed to be KREEP-like or flatter, despite the extreme La/Lu fractionation manifested across the spectrum of SNC compositions.

Also problematic is the high proportion (probably $\gg 10$ wt%, making realistic assumptions concerning Sr) of the enriched component that must be assimilated into the HD starting material to reach the average SD composition. Alternatively, the model may be over-simplified; MD and SD shergottites are not necessarily simple dilutions of HD matter by a single, uniform “crust” component.

The notion of a highly depleted source region, at least for HD shergottites, and mixing with less depleted crustal matter to account for the slightly depleted (SD) shergottites like Shergotty and Zagami, goes back at least to Nyquist et al. (2001b), and in some respects to Borg et al. (1997). Recent studies of shergottites (Wadhwa, 2001; Herd et al. 2001, 2002; Goodrich et al. 2003; Herd, 2003) have added an important new constraint by documenting a correlation between geochemical depletion/enrichment and oxygen fugacity (Fig. 15). For La/Lu, this correlation extends to the nakhlites and Chassigny. Again, despite their cumulate nature, the nakhlites and Chassigny have REE patterns (including La/Lu) lower but roughly parallel to the patterns of their parent melts (see above, and cf. Wadhwa and Crozaz, 1995). The quantitative $f_O^2$ data shown for nakhlites are mainly from a recent abstract (Syzmanski et al. 2003), but Wadhwa and Grove (2002 report that the Gd/Eu technique yields similarly high typical $f_O^2$ for an unspecified set of nakhlites. Chassigny’s $f_O^2$ is not yet well constrained. The $f_O^2$ plotted in Fig. 15 (QFM+0.5) is based on the spinel technique of Wood (1991; cf. Goodrich, 2003, Herd, 2003) and our own new Cr-spinel analyses; but Chassigny’s spinels may not be entirely equilibrated (Floran et al. 1978), so this $f_O^2$ is only a highly approximate result, assuming a relatively high “equilibration” T of 1000°C.

It is generally assumed that the shergottites are a representative rock type in relation to the upper mantle melts (e.g. Wadhwa, 2001; Herd, 2003). If the relationship between depletion and $f_O^2$...
manifested among the shergottites (Fig. 15) reflects a general feature of the martian mantle-crust system, and if the spinel-derived (e.g. Herd, 2003) and crystallization simulation-derived (McKay et al. 2002, 2004) $f_{O_2}$ results are correct (Wadhwa’s are systematically lower by ~ 2 log units), then most of the martian upper mantle is at ~ QFM –4 and also highly depleted.

Herd (2003) suggested that the diversity in both depletion and $f_{O_2}$ among the shergottites (Fig. 5, 15) may be after-effects of a magma ocean episode. He argued that rather than reflecting mixing between primitive, reduced basaltic magmas and oxidized crust (Nyquist et al. 2001b), the depletion diversity among shergottites mainly reflects heterogeneity within the mantle, dating back to magma ocean crystallization. Herd (2003) assumes that the oxidizing component (basically Fe$^{3+}$) was probably closely linked with H$_2$O, and suggests H$_2$O may have originally concentrated together with trapped melt during crystallization of the magma ocean. Herd (2003) and Treiman (2003) also note that the H$_2$O-rich component in the mantle would likely become remobilised and cause complex, heterogeneous metasomatic alteration of the upper mantle. These suggestions followed a brief argument by Wadhwa and Grove (2002) that the mantle $f_{O_2}$ and depletion heterogeneities formed during early differentiation because both depletion and loss of volatiles “such as H$_2$O and CO$_2$” affected most of the mantle, but shallow regions beneath ancient cratons were exceptionally stable (against convective stirring), and these subcratonic reservoirs were subsequently “variably metasomatized and oxidized by the melts and degassed volatiles from the deep mantle.”

The nature of the source region for the nakhlites is not certain but it is distinct from those of the shergottites. Jones (2003) suggests that the nakhlite source region represents the lower mantle; and that it stabilized below the shergottite source region because it is less depleted, and thus richer in Al$_2$O$_3$, which at mantle pressures forms dense garnet. From a physical standpoint, this seems a plausible model. Besides having more majorite, the fertile mantle would presumably be comparatively FeO-rich. Indeed, the low Mg# of nakhlites in comparison to similarly mafic-concentrated (i.e. Al$_2$O$_3$-poor) cumulate shergottites (Fig. 16) suggests a source region considerably lower in Mg# than the shergottite source. Density constraints reviewed by Elkins-Tanton et al. (2003) indicate that for a given Mg#, majorite is ~0.26 g cm$^{-3}$ denser than olivine (and low-Ca pyroxene; however, majorite is ~0.08 g cm$^{-3}$ less dense than $\gamma$-olivine, the stable olivine in the ~18 vol% of the martian mantle that is below ~1300 km). A diminution in Mg# of 10 mol% (as an example) translates into higher density by ~ 0.12 g cm$^{-3}$, so the Mg# effect might be even more important than a moderately higher majorite proportion in
impacting negative buoyancy to a nakhlite source. An advantage of this model is that the source is implied to be garnet-rich, and retention of even minor garnet in a restite should impart a high La/Lu to the escaping melt; yet nakhlites (and their parent melts) have La/Lu much greater than chondritic (Fig. 9).

Magma Ocean and spectroscopic signatures of large-scale crustal differentiation

If a martian magma ocean (Borg and Draper, 2003; Elkins-Tanton et al., 2003; Herd, 2003; Jones, 2003) existed, its end-product crust might be expected to bear some resemblance to the highly anorthositic (Al$_2$O$_3$-rich) crust that is often assumed to have formed by magma ocean plagioclase flotation on the Earth’s Moon. Over most of the Moon’s surface, soils have ~25-28 wt% Al$_2$O$_3$ (Warren, 2004), a proportion far higher than what could plausibly form by alternative models (i.e. “serial magmatic” piecemeal emplacement of mantle-derived partial melts). However, soils at four widely separated martian sites have only about 10 wt% Al$_2$O$_3$ (Foley et al. 2003; Rieder et al. 2004).

Data from orbital thermal emission spectrometry (TES, Mars Global Surveyor: Bandfield et al. 2000) have been interpreted as two major compositional terranes: Type 1, basaltic (generally in the southern, older highlands) and Type 2, andesitic (generally in the northern, relatively young, low-lying plains). However, in an alternative interpretation of TES data, Wyatt et al. (2004) noted that there was a correlation between the Type 2 terranes and near-surface ice at middle to high latitudes. They suggested that basalt, weathered by limited interaction with H$_2$O over short periods of time, could equally provide the fit rather than andesite for TES Type 2 surfaces. The geochemical evidence for basaltic andesites has been discussed previously; the modelled mineral abundances within the basaltic and andesitic spectroscopic signatures are discussed in a later section.

Orbital γ-ray spectrometry, a technique that “sees” beneath the surface soil layer (by roughly 1 m), shows higher FeO (14 to 19%) in the northern lowlands than in the highlands (10 to 14%) (Boynton et al. 2004). Thus, the γ-ray data do not support the interpretation (Bandfield et al. 2000) that the northern crust is generally andesitic (andesites on the less FeO-rich Earth average about 7 wt% Fe: Chayes, 1969). However, other aspects of the γ-ray spectrometry data – K abundances – do support the idea of andesitic material in the northern lowlands. Summing up, the spectroscopic (γ-ray and TES) evidence bearing on andesites or basaltic andesites on Mars is inconclusive. Neither the spectroscopic data nor in situ soil analyses provide evidence for the preservation of early anorthositic crust.
The recent literature contains many suggestions of an early martian magma ocean (e.g. Borg and Draper, 2003; Elkins-Tanton et al., 2003; Herd, 2003), but these models seldom mention the obvious possibility that a magma ocean would have engendered an anorthositic flotation crust, as apparently happened on the Moon. Any early crust might conceivably have been recycled into the mantle, but flotation anorthosite is especially buoyant, and major involvement of surface plates in mantle convection is an unlikely scenario for Mars (e.g. Warren, 1993). Borg and Draper (2003) have proposed models of martian magma ocean crystallization in which plagioclase flotation is a non-issue, because no plagioclase ever forms. Borg and Draper (2003) assumed that crystallization of pressure-stabilized majorite (garnet) and clinopyroxene so depleted the melt’s Al$_2$O$_3$ that it never reached saturation with plagioclase. However, based on a wealth of experimental constraints (e.g. Herzberg and Zhang, 1996; Bertka and Fei, 1997), igneous majorite crystallization at pressure less than (conservatively) 1 GPa is impossible. Realistically, even assuming closed-system fractional crystallization at the high pressures associated with the base of the magma ocean, enough Al$_2$O$_3$ remains in the melt to eventually form many tens of kilometers (equivalent thickness) of plagioclase (Warren, 1989). Furthermore, the late-stage magma ocean (a) is prone to crystallize at its top as well as along its base (e.g. Morse, 1982); and (b) is more realistically only the upper, fully molten subvolume of a magmasphere; i.e. the melt zone is probably an open system that gets replenished with fresh (and compositionally primitive) melt ascending out of the deeper interior, even during what is mainly a process of crystallization (e.g. Warren, 1989).

A key parameter during crustal genesis by magma ocean plagioclase flotation is the density of the underlying melt. The melt will be saturated in pyroxene (and probably also olivine) as well as plagioclase, and the nascent crust will tend to entrain a proportion of these mafics (or melt that promptly crystallizes mafics) sufficient to impart an approximately neutral density. Warren (1990), considering a lunar magma ocean, showed that the relevant density-balance equation is:

$$v_{\text{plag}} = \frac{\rho_{\text{maf}} - \rho_{\text{melt}}}{\rho_{\text{melt}} - \rho_{\text{plag}}}$$  \[3\]

where $v_{\text{plag}}$ is the volume fraction of plagioclase in the nascent crust, $\rho_{\text{maf}}$ is the average density of the mafic component of the nascent crust, $\rho_{\text{melt}}$ is the density of the melt, and $\rho_{\text{plag}}$ is the density of the nascent plagioclase. The relevant temperature here is roughly 1200°C, and the relevant pressure is low (even at the base of the final lunar or martian crust, only about 0.3-0.6 GPa). The low-pressure density of a silicate melt can be calculated to good precision based on its composition and the partial molar
volumes of its constituent oxides (Lange and Carmichael, 1990). The quantity $\rho_{\text{plag}}$ is nearly a constant; it is slightly (0.03 g cm$^{-3}$) lower for a martian composition as opposed to a Na-poor lunar one. By far the most important determinant of $\rho_{\text{melt}}$, for the range of conceivable melt compositions, is FeO. The value of $\rho_{\text{melt}}$ (assuming analogous varieties of mafic silicate) is also basically determined by FeO, and so is almost directly proportional to $\rho_{\text{melt}}$. Thus, to a first approximation, the volume fraction of plagioclase in the nascent crust $v_{\text{plag}}$ is determined by one major variable, $\rho_{\text{melt}}$, which is in turn very largely a function of the melt FeO concentration.

The martian primitive mantle composition (constrained by geophysics as well as petrology) is fully twice as FeO-rich as the Earth’s mantle (Bertka and Fei, 1998; Dreibus and Wänke, 1985). A disparity in Mg# between martian rocks and analogous terrestrial mafic-ultramafic igneous rocks is obvious (Fig. 16). The martian mantle is probably also ferroan in comparison with the Moon, judging from the disparate trends for cumulate/phyric samples (filled symbols) on Fig. 16. Lunar regolith compositions also extrapolate to a high, Earth-like Mg# (Warren, 2004). It seems likely that FeO is higher by roughly a factor of 2 in Mars’ crust-mantle system than in the Moon.

The relationship between initial melt FeO content and melt FeO (or $\rho_{\text{melt}}$) near the end of a magma ocean crystallization sequence is far more complicated than a direct proportionality. However, in the case of the lunar magma ocean, the value of $\rho_{\text{melt}}$ can be estimated based on the observed consistently low-Mg# and high-$v_{\text{plag}}$ nature of the distinctive “ferroan anorthosites” that are the only plausible representatives of a plagioclase-flotation crust (e.g. Warren, 2004). The implied $\rho_{\text{melt}}$ is roughly 2.76-2.85 g cm$^{-3}$, depending on whether assessed mainly based upon the observed high $v_{\text{plag}}$ (roughly 93 vol%) implied by the ferroan anorthosites, or mainly upon forward modelling of magma ocean crystallization (Warren, 1990). KREEP-basaltic melts (Ryder, 1985; Salpas et al. 1987) are even less dense (2.71-2.76 g cm$^{-3}$). In the case of Mars, one approach to estimating $\rho_{\text{melt}}$ is to assume the late-stage magma ocean compositionally resembled the average of the four shergottites at or near plagioclase saturation (Fig. 17); which implies $\sim$2.81 g cm$^{-3}$ (at 1200°C, assuming $<<$1% H$_2$O). However, two of these shergottites are depleted (EET79001-B is MD, QUE94201 is HD), and in a very general way incompatible element depletion should correlate with FeO/MgO (i.e. melt density) depletion. Forward modelling, using assumptions analogous to those that imply $\sim$2.85 g cm$^{-3}$ for the late lunar magma ocean, suggests a late-stage $\rho_{\text{melt}}$ of roughly 2.85-2.94 g cm$^{-3}$ (Warren, 1989).
Fig. 17 shows these density estimates plotted vs. the $v_{\text{plag}}$ implied by eqn. [3], assuming a range of plausible $\rho_{\text{mof}}$. It would be unrealistic to expect such estimates to be accurate to much better than 0.2 g cm$^{-3}$. The only reasonably confident inference is that a late-stage martian magma ocean was probably denser than its lunar counterpart, by very roughly 0.1 g cm$^{-3}$. But even this is an important inference. The implied $v_{\text{plag}}$ is not 81-93 vol%, as in the lunar scenario, but 67-85 vol%. Even after magma ocean solidification, the same basic constraints apply: A martian crust with 76 vol% plagioclase (the nominal, mid-range result in Fig. 17) is about equally as buoyant over the dense, FeO-rich martian mantle as a lunar crust with about half as much mafic silicate (87 vol% plag) over the less dense, relatively FeO-poor lunar mantle. This moderation of the flotation crust’s composition may be part of the explanation for the lack of spectroscopic evidence for an anorthositic crustal remnant on Mars.

The existence of a magma ocean during Mars’ history might account for some of the compositional features of the SNCs such as Al-depletion (Al being trapped in plagioclase of the magma ocean). The $f_\text{O}_2$ and LREE enrichments sometimes thought to be a crustal contaminant component in the shergottites might instead have resulted from magma ocean differentiation followed by a mobilisation of H$_2$O and Fe$^{3+}$ (Herd, 2003). An alternative crustal origin for the geochemical diversity among SNCs is discussed further in relation to the source-crater locations of the SNCs.

**Crystallization and Ejection Ages of the SNCs**

The ages at which the SNC meteorites crystallised on Mars have been determined in a variety of ways including Rb-Sr, Sm-Nd and $^{39}$Ar-$^{40}$Ar and this - together with Cosmic Ray Exposure (CRE) ages - are reviewed in Nyquist et al. (2001a). In summary, the crystallization ages fall within 5 groups. ALH84001 is the oldest at 4.5 Ga, Chassigny is 1.3 Ga, the nakhlites 1.3 Ga, peridotites 180 Ma and basaltic shergottites 165-475 Ma. From these ages it has been taken that ALH84001 is a fragment of Noachian crust and so probably from the southern highlands. The other 29 meteorites date from the Amazonian System, which is approximately <2.9 Ga (Hartmann and Neukum, 2000). These young ages of the shergottites suggest that they were derived by impacts onto the northern hemisphere and the 2 main likely source regions are Tharsis and Elysium-Amazonis.

The closeness of the Chassigny and nakhlite crystallization ages, which overlap within the errors, might mean that they are comagmatic although no firmly established petrogenetic model to explain this exists. Their parental melts are both FeO-rich, Al$_2$O$_3$-poor, LREE>HREE, and basaltic (Johnson et al.
Further evidence for a link between Chassigny and the nakhlites is that they both have CRE ages of 11 Ma. The CRE ages of the SNC meteorites fall within up to 7 groups at 20, 15, 11, 4.5, 3, 1.3 and 0.7 Ma, leaving the possibility of 7 ejection events from Mars between 20 and 0.7 Ma (Nyquist et al. 2001a). There is a clustering of ejection ages around 3 Ma for 4 of the basaltic shergottites (Los Angeles, QUE94201, Shergotty and Zagami). The peridotitic shergottites may not have been ejected in the same event e.g. LEW88516 and Y793605 were apparently ejected at 3.9 ± 0.4 Ma at 4.7 ± 0.5 Ma but the close petrographic and geochemical similarities between the peridotitic shergottites may indicate another explanation. ALH84001 has a calculated ejection age of 15 Ma.

All but one of the possible ejection events were from the northern lowlands or Tharsis. However the ancient highlands occupy about 60% of the martian surface. Furthermore, only ~ 12% of the martian surface is composed of lavas young enough to be associated with the shergottites (Tanaka et al. 1992; Keszthelyi et al. 2000). The reason for this over representation of young, shergottitic material from the northern lowlands is not certain. However, it is possible that some secondary collisions in space have acted to skew the statistics and exaggerate the number of impacts on Mars that are being sampled (Nyquist et al. 2001a). Nyquist et al. and Warren and Kallemeyn (1997) also noted that uncertainty in production rates mean that the distinct possibility exists, for instance, that all of the peridotitic shergottites were ejected in the same event from one crater.

Therefore the number of ejection events that have currently been sampled remains uncertain although 4 is a reasonable minimum estimate. Current modelling suggests that craters ≥3 km (Head and Melosh, 2000) can be associated with ejection of material from Mars at the escape velocity of 5 kms⁻¹.

**Source regions and the martian surface**

The young SNC ages demonstrate the existence of igneous activity on the martian surface to relatively recent times. Although as noted previously the high proportion of young shergottites is hard to reconcile with the martian surface there is increasing evidence – particularly from the Mars Global Surveyor camera – for young lava flows on the surface of Mars. Hartmann and Neukum (2001) conducted impact crater counts on some flows within Elysium Planitia and suggested ages of 10 Ma or less. Young lava flows (<500 Ma) cover approximately 12% of the volcanic surface area of Mars (Keszthelyi et al. 2000). The potential SNC source areas in Tharsis and Elysium-Amazonis Planitia
both contain lava flows thought to be <100 Ma (Hartmann and Neukum, 2001). Some of the basaltic flows on Mars may be up to 1-2 km thick and extend over 100s km² consistent with an Fe-rich, low viscosity nature.

Mouginis-Mark & Yoshioka (1998) studied the distribution and morphology of 59 lava flows in Elysium Planitia. They measured average widths from 3 to 16 km, with many of the flows exceeding 100 km in length. Photoclinometry was used to derive an average thickness of 40-60 m for these flows, indicating that individual flow volumes range from 18 to 70 km³. Thus formation of nakhlite/Chassigny and perhaps the final crystallisation of basaltic shergottites can potentially be accommodated within such lava flows. Peridotitic shergottites may have crystallised in situ as plutonic or intrusive material in deeper crustal levels.

Hamilton et al. (2003) reported the results of a search using TES data for spectral signatures close to those of the SNC types. Although some olivine and orthopyroxene-rich areas were identified, no fits to the laboratory SNC spectra were found. However, nakhlite-like materials were tentatively identified at near detection limits in the Valles Marineris area.

Bandfield et al. 2000 highlighted areas e.g. Chryse and Acidalia Planitia where what they suggested was an andesitic component (Type 2 surface) was seen. The Elysium-Amazonis and Tharsis regions do not have high detected proportions of the Type 1 or 2 end-member TES spectral signatures because the presence of surface dust preclude any useful spectra. The ancient highlands are interpreted to have a mainly basaltic signature although some areas within it (including possible source regions in Valles Marineris for the rocks on the Chryse flood plain surface) also have andesitic signatures. The spectra for the ‘basaltic’ signature was plagioclase 50%, augite 5%, sheet silicate 15%; for the ‘andesitic’ component the fit was plagioclase 35%, augite 10%, sheet silicate 15%, K-rich glass 25%. The basaltic and andesitic-like signatures dominate the surface of Mars with only relatively minor olivine-rich or haematite-rich areas. However, Hamilton et al. (2000, 2003) and Bibring and Erard (2001) noted that the basaltic TES spectra do not closely match those of the shergottites. Furthermore, Wyatt & McSween (2002) and Wyatt et al. (2004) modified the deconvolution mineralogy, significantly decreasing the estimated feldspar in both terrain types.

The calculated water contents of melt inclusions within shergottites suggest that substantial volumes of water were brought to the martian surface during the ascent and eruption of basaltic magmas. For instance, taking 1.8 wt% for the water content of an erupting basalt, then for a lava flow
of 70 km$^3$, and a magma density of 2.85 g cm$^{-3}$, 3.6 x 10$^6$ m$^3$ of water would be released. Scott et al. (1998) calculated the volume of individual Tharsis shield volcanoes, including their roots, as 1.6 x 10$^{16}$ m$^3$. This could have produced 8.2 x 10$^{10}$ m$^3$ of water which is ~7% of the volume of the maximum north polar cap (Zuber et al. 1998).

We have demonstrated that conditions may have been right for the existence of a magma ocean during Mars’ history. Some researchers have suggested that this magma ocean caused the geochemical heterogeneities present in the SNCs, in particular the link between LREE contents and $f$O$_2$ (Herd, 2003). As described above we suggest that alternative models (e.g. Nyquist et al. 2001b) based on crustal contamination do not fully account for the range of isotopic and chemical abundances present in the SNCs.
**Synthesis and Conclusions**

The most abundant group of SNC meteorites are called the basaltic shergottites. Their composition is similar to that of rocks analysed at the *Opportunity* landing site and the basaltic component that comprises much of the martian surface regolith and underlying geology in terms of Fe-enrichment. However the *Spirit* rocks are picritic and also more alkali-rich than the basaltic shergottites. Olivine-phyric shergottites form another recognizable group of the shergottites which accumulated phenocryst or xenocryst olivine grains from a separate olivine-saturated basaltic melt. These large olivine grains were not derived by disruption of peridotite shergottite sources because the olivine-phyric shergottites generally have a more highly depleted geochemistry than the peridotitic shergottites. The 6 peridotite shergottites have the clearest cumulate textures of the SNCs and differ from the other shergottites in their low proportion of feldspatic material and high proportions of olivine. The cores of pyroxene in basaltic shergottites crystallised slowly at depth from melts which at least in some cases were H$_2$O-bearing (e.g. 4 - 6 km, ≤1.8 wt% H$_2$O) followed by more rapid crystallisation of the Fe-rich rims in a near-surface intrusive or extrusive setting.

The 6 ol-pyroxenite nakhlites formed as cumulates in a thick e.g. ~100m lava flow from the accumulation of augite followed by olivine. Trapping of varying amounts (5-20%) of basaltic, interstitial melt – the nakhlites in the upper parts of the parental lava flow having the higher proportions of trapped melt - has given the nakhlites their LREE-enriched geochemical signature. The dunite Chassigny which has near identical ejection and crystallisation ages to the 6 nakhlites may also be associated with them.

The martian mantle source region has over twice the FeO contents of the terrestrial mantle and the SNC compositions reflect this in their Fe-enrichment compared to analogous terrestrial and lunar rocks. Another compositional feature of the SNCs is their low Al contents which reflect depletion of source regions, perhaps due to the formation of a magma ocean. However discordance between Mg# and Al$_2$O$_3$ contents of the nakhlite and other SNC groups shows that the SNC melts were derived from mantle source regions with differing depletion histories. The modelled martian magma ocean would have a lower proportion of plagioclase and lower density than the lunar one. Inferred noble metal contents in the martian mantle calculated from the SNCs suggest that – like the Earth – Mars underwent a later accretion of chondritic material.
In addition to petrographic classification, we suggest that the basaltic, olivine-phyric and peridotitic shergottites can also be subclassified geochemically on the basis of their LREE-depletion into HD (highly depleted), MD (moderately depleted) and SD (slightly depleted). The SD shergottites mainly correspond to the basaltic shergottites, the MD to peridotitic shergottites and the HD correspond to olivine-phyric shergottites. However LD, MD and SD all have some members from other petrologic groups. This is an important classification because the La/Lu ratios (i.e. depletion) of the shergottites correlate with $\log_{10}f_O$. The LD shergottites and also the nakhlites and Chassigny have $\Delta$QFM of -1 to +0.5 and the HD shergottites have $\Delta$QFM of -2.5 to -3.5. From these results it is inferred that the most reduced martian mantle has an oxygen fugacity of QFM –4.

The melt compositions of the SNCs – either known directly from the whole rock composition or calculated from melt inclusions within cumulate phases – do not show any clear evidence for an andesitic component. The best data available from the Pathfinder rock analyses suggests those rocks may be basaltic andesites i.e. with slightly higher SiO$_2$ and Na$_2$O + K$_2$O than the basaltic shergottites (although it is possible that this chemistry reflects contamination of the Pathfinder analyses by alteration rinds). Spectroscopic data in support of the geochemical evidence for an andesitic component in the northern lowlands of Mars is not yet conclusive. However, the existence of a K- and Fe-enriched component in parts the northern lowlands distinct from the basaltic signature in the remaining northern lowlands and southern highlands is established from TES and $\gamma$-ray spectroscopy. The formation mechanism for such large scale magmatic heterogeneities is not clear but might involve fractionation of basaltic magmas trapped in magma ocean rocks or the fractionation of shergottitic compositions under hydrous conditions.

The absence of an andesitic-like chemical signature in the SNCs suggests that they were derived from areas in the northern lowlands where the K-rich ‘andesitic’ spectral signature is absent or in low abundance. Two likely regions are the Tharsis region of shield volcanoes and the Elysium-Amazonis volcanic plains. These regions also contain young volcanic rocks compatible with the relatively young ages of the SNCs. The crystallization ages fall within 5 groups. ALH84001 is the oldest at 4.5 Ga, the nakhlites and Chassigny 1.3 Ga, peridotitic shergottites 180 Ma and basaltic shergottites 165-475 Ma. On the basis of these ages it is clear that only the ALH84001 orthopyroxenite is derived from the ancient highlands: this also shows the sampling bias in our current collection of 31 SNCs because ~60% of the martian surface is comprised of the ancient highlands. The SNCs were ejected from Mars.
in between 4 and 7 impact events but uncertainties in the calculation of ejection ages means that the grouping of samples with ejection events is not always clear. However the ejection of the Nakhliites and Chassigny in one event 11 Ma is well established.

The correlation of La/Lu with $\log_{10} f_O$ in the SNCs may be due to magma ocean-related mantle heterogeneities involved with the mobilisation of H$_2$O and volatiles. We favour this explanation (i.e. mantle heterogeneity) over alternative explanations that the correlation is due to variable contamination with an oxidised, Sr-rich, high La/Lu hydrous fluid in crustal magma chambers. We find that mixing calculations (La/Lu vs. $^{87}\text{Sr}/^{86}\text{Sr}$) between inferred crustal components and HD shergottite samples do not fully satisfy the range of isotopic and geochemical abundance patterns present in the SNCs.

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References


**Figures**

Fig. 1. Shergotty basaltic shergottite in thin section. px clinopyroxenes (pigeonite and augite grains), mk maskelynite, mt magnetite. The lighter margins of the pyroxene grains are Fe-enriched rims, and this is typical of both basaltic and olivine-phyric shergottites. Black parts are holes in the section. Back-scattered electron image. Field of view 5 mm.

Fig. 2. Pyroxene (augite and pigeonite) compositions in basaltic shergottites and 2 olivine-phyric shergottites (SaU005 and DaG476) in dashed line areas. The pyroxenes are all notably Fe-enriched,
with the most extreme enrichment seen in Los Angeles. The olivine-phyric shergottites have the least Fe-rich augite and pigeonite. Analyses made with Cameca SX50, Natural History Museum.

Fig. 3. Y000749 nakhlite (a meteorite paired with Y000593) in thin section. Radiating laths of plagioclase are seen in the mesostasis. Planar fractures, running NE-SW, are present in the marked, large olivine grain in the centre-right of the image, and these are a feature of shock metamorphism. ol olivine, px pyroxene (augite), mes mesostasis, mt Ti-magnetite, plane polarised view, 1000 µm scale.
Fig. 4. Pyroxene and olivine compositions in the nakhlites. The main mineral phase in the nakhlites is an Fe-rich augite (En$_{0.37-0.62}$Wo$_{0.37-0.43}$Fs$_{0.24-0.41}$) with lesser amounts of pigeonite (En$_{48-54}$Wo$_{6-13}$Fs$_{35-39}$). There is little discernible compositional variation between 4 of the nakhlites (Nakhla, Lafayette, Governador Valadares, NWA998). However, Y000593 and NWA817 have the most Fe-rich augite core compositions – extending to Fs$_{0.4}$ indicating that they have more fractionated assemblages. The arrow indicates the zonation of the augite rims to towards hedenbergite-like compositions, and this is most extreme in NWA817 augite. NWA817 also has the greatest range of olivine compositions (Fo$_{14-44}$) suggesting that it has undergone the least equilibration. Olivine grains compositions are plotted beneath the pyroxene quadrilateral. Their Mg# ratios are not in equilibrium with the pyroxene compositions or with the bulk Mg# of Nakhla (also plotted, 0.5) and this feature may be due to late crystallization of the olivine relative to the augite cores (e.g. see Fig. 9). The pyroxene and olivine compositions are from this study apart from the Y000593 olivine (Mikouchi et al. 2003) and the NWA817 augite, rims and olivine data (Sautter et al. 2002).
Fig. 5. CI-chondrite-normalized rare-earth element (REE) patterns for basaltic shergottites. Data plotted are from the compilation of Meyer (2003) and Table 5. Patterns of the highly depleted (HD) type are extremely depleted in light REE, especially La, despite being nearly flat among the heavier REE (Tb-Lu). Patterns of the slightly depleted (SD) type are nearly flat but slightly bow-shaped, with La/Tb and Lu/Tb both ~ 0.7 × CI. The moderately depleted (MD) type is intermediate.
Fig. 6. Martian basalt and peridotitic shergottites compositions (wt%). A. CaO v. MgO; B. Al$_2$O$_3$ v. MgO; C. TiO$_2$ v. MgO. The basaltic shergottites and *Pathfinder* analyses can be seen to have distinct compositions, *Pathfinder* analyses having lower, CaO, TiO$_2$ and higher Al$_2$O$_3$. Also plotted are compositional fields (dashed curves) for terrestrial oceanic basalts (Basaltic Volcanism Study Project, 1981). These show the generally low Al$_2$O$_3$ contents and similar CaO of most SNCs compared to terrestrial basalts. The Gusev analyses are distinct from both the basaltic and olivine-shergottites,
having higher MgO, lower TiO₂ and CaO than the basaltic shergottites but lower MgO and higher Al₂O₃ than the olivine-shergottites. See Table 4. for data sources.

Fig. 7. Martian basalt compositions, Na₂O + K₂O v. SiO₂ (wt%). The *Pathfinder* rock and soil-free rock analyses from Wänke et al. (2001) and Foley et al. (2003) are plotted. The *Spirit* analyses (McSween et al. 2004) are similar to picritic basalts, whereas the *Opportunity* analysis has a higher SiO₂ content. Neither the *Spirit* nor the *Opportunity* analyses have similar compositions to the *Pathfinder* analyses. The latter have alkali-, silica-rich basaltic-andesite to andesite compositions. Ol-Shergottites are olivine-phyric shergottites; B. shergottites are the basaltic shergottites. Compositional fields from Cox et al. (1979).
Fig. 8. Martian rock bulk compositions on the olivine-plagioclase-silica phase diagram of Longhi (Longhi and Pan, 1988; Longhi, 1991), based on oxygen (~volume) units, projected from wollastonite. Conversion from molar into oxygen units is done by dividing the molar percentages by the number of oxygens in the molecular formula (4 for olivine, 8 for plagioclase, 2 for silica) and renormalizing to 100%. Boundaries shown in black are appropriate for melts with Mg# ~40-50 mol% and Na/(Na+Ca+K) ~20 mol%; boundaries shown by grey lines are for higher Na/(Na+Ca+K), ~40 mol% (see Longhi and Pan, 1988 and 1991). Meteorite compositions are based on literature data compiled in Meyer (2003), augmented by Dreibus and Jagoutz (2004) and Table 5 data of DaG476/735, SaU005, Y000593 and Y000749. In order of increasing “Pl,” the plotted individual basaltic shergottites (blue symbols) are SaU005, DaG476, Y980459, Dhofar 019, Zagami, NWA480, Shergotty, QUE94201, Dhofar 378 and Los Angeles. Open symbols denote samples whose bulk compositions are believed to closely resemble their parent melts. Also shown (open symbols connected to meteorites by dotted lines) are modelled parent melts for Nakhla (Treiman, 1993), the LEW88516 peridotitic shergottite (Harvey...
et al. 1993), SaU005 (Goodrich, 2003) and Shergotty (Hale et al. 1999). Red crosses denote compositions obtained in situ by the *Pathfinder*, *Spirit* and *Opportunity* probes. The plotted Pathfinder compositions are the averages of 5 analyses (each) reported by Wänke et al. (2001) and Foley et al. (2003); as indicated in Table 2, these teams derived disparate results, particularly for SiO₂.

![X-ray maps](image)

**Fig. 9.** X-ray maps (A. is Fe Kα and B. is Mg Kα) of nakhlite (Governador Valadares) cumulate texture. The early-formed augite cores, with homogenous compositions are seen together with their Fe-rich and Mg-poor rims. The large olivine grain at the bottom of the figure has crystallised after the augite cores but before their Fe-rich rims. Scale bars 200 μm.
Fig. 10. REE abundances of whole Nakhla and Y000593 (Table 6) and NWA817 (Sautter et al. 2002) samples together with fractional crystallization model. In this model primitive ol-augite cumulates are formed from a fractionated basaltic melt. The degree of fractional crystallization of this melt $F$ (i.e. fraction of melt remaining) = 0.6. The REE abundances of this melt and that of the cumulates are shown. Addition of 5-20% of this melt to the cumulate, corresponding to trapping of interstitial melt in the nakhlites’ mesostases, produces calculated REE abundance similar to those of the Nakhla, Y000593 and NWA817 whole samples. See text for details of the model. REE are normalised to CI abundances.
Fig. 11. Trace element abundances of nakhlite: Nakhla, Lafayette, Governador Valadares, Y000593, NWA998 minerals. Open symbols are cumulus augite, corresponding closed symbols are mesostasis (formed from trapped melt). The augite trace element abundances within the 5 nakhlites are indistinguishable. See Table 6 for representative data. Dashed line is Nakhla parent melt of Wadhwa and Crozaz (1995). Normalised to CI chondrites.
Fig. 12. Schematic diagram of nakhlite cumulate pile. The overall thickness e.g. in a large lava flow is ~100-200 m. NWA817 has the highest proportion of trapped, interstitial melt (light blue). This is reflected in its higher REE abundances (Fig. 9). Lafayette may have crystallised in the lowest part of the cumulate pile, as reflected by its equilibrated mineral compositions and lack of mineral zonation. Some of the augite grains show twinning on (100). Diagram after Mikouchi et al. (2003).
Fig. 13. Correlations between the highly siderophile element Ir and MgO for mafic igneous rocks from Mars, Earth, and the Moon. For the martian HD olivine-phyric shergottite SaU005, the plotted Ir value is the 2:1 weighted average of a direct measurement (13 ng/g: UCLA, Table 5) and an extrapolation from a vastly lower Os result (0.62 ng/g) found by Walker et al. (2002). The datum plotted for lunar sample 15426 (green mare-pyroclastic glass) is an upper limit (Walker et al. 2004). Other sources for martian data are as compiled by Meyer (2003), plus Jones et al. (2003; excluding suspiciously high data — we suspect contamination — for ALH84001 and three shergottitic basalts) and Table 5 for the DaG735 pair of DaG476. Terrestrial and lunar background data are from sources compiled in Warren et al. (1999), most notably, for the MgO-rich lunar samples 67667 and 72415, Warren and Wasson (1979), Ebihara et al. (1992) and Morgan and Wandless (1988).
Fig. 14. Shergottites show a strong correlation, and yet a clustering into three subgroups, on a plot of La/Lu vs. initial $^{87}\text{Sr}/^{86}\text{Sr}$ (normalized to a typical shergottite age of 175 Ma). Symbols are analogous to those in Fig. 8. Nakhlites and Chassigny (and to a lesser extent the ancient orthopyroxenite ALH84001) plot well off the shergottite trend. Literature data used for this plot are from the same array of sources as for Fig. 8. Also shown are mixing parabolas derived from four models that are discussed in the text.
Fig. 15. Oxygen fugacity correlates with the ratio La/Lu among nearly all martian meteorites, ancient orthopyroxenite ALH84001 being a lone outlier. Symbols are analogous to those in Fig. 8. Literature $fO_2$ data used for this plot are mainly based on opaque mineral analyses: mainly from Goodrich (2003), Herd (2003) and Syzmanski et al. (2003); also Cahill et al. (2002), Warren et al. (2004), and unpublished Open University/Natural History Museum data (Table 3) for Chassigny. Caveat: $fO_2$ results for Chassigny and ALH84001 (Herd and Papike, 1999) are only approximate. Important additional constraints on $fO_2$ for basaltic shergottites are available from crystallization experiments (McKay et al. 2002, 2004), and from the Gd/Eu technique of Wadhwa (2001); the Gd/Eu results tend to confirm the correlation with La/Lu, but are systematically offset from spinel-based results by about -2 log units; the technique is still undergoing calibration (McCanta et al. 2002). Data for La/Lu are from the same combination of sources as for Fig. 8.
Fig. 16. Martian meteorites show systematically low Mg# when compared with analogously mafic, Al$_2$O$_3$-poor Earth and Moon rocks. Symbols for martian samples are analogous to those in Fig. 8; the single Pathfinder point is the average of the Wänke et al. (2001) and the Foley et al. (2003) versions of the average for five rocks. Data for martian meteorites are from the same sources as for Fig. 8. Data for komatiites and komatiitic basalts are from BVSP (1981). Data for lunar samples are from the compilation of Warren (2004), augmented with data for lunar mare-pyroclastic glasses from Shearer and Papke (1993). The many lunar mare basalt (and pyroclastic glass) compositions shown are averages of various discrete varieties (not necessarily representing individual lavas; e.g., all Apollo 12 basalts are averaged together). The two KREEP basalts shown are individual rocks (15386 and 72275; compositions from Ryder, 1985 and Salpas et al. 1987).
Fig. 17. Volume fraction of plagioclase $v_{\text{plag}}$ in a magma ocean flotation crust, calculated as a function of melt density using eqn. [3]. The high melt density expected for a late-stage martian magma ocean (see text) probably translated into a relatively moderate $v_{\text{plag}}$. 

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