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142Nd Evidence for Early (>4.53 Ga) Global Differentiation of the Silicate Earth

M. Boyet* and R. W. Carlson

New high-precision samarium-neodymium isotopic data for chondritic meteorites show that their 142Nd/144Nd ratio is 20 parts per million lower than that of most terrestrial rocks. This difference indicates that most (70 to 95%) of Earth’s mantle is compositionally similar to the incompatible element–depleted source of mid-ocean ridge basalts, possibly as a result of a global differentiation 4.53 billion years ago (Ga), within 30 million years of Earth’s formation. The complementary enriched reservoir has never been sampled and is probably located at the base of the mantle. These data influence models of Earth’s compositional structure and require revision of the timing of global differentiation on Earth’s Moon and Mars.

Radiogenic isotope tracers are often used to understand the chemical evolution of planetary bodies. In one system, Sm decays to Nd via two radioactive decay schemes: 146Sm–142Nd [half-life T(1/2) = 103 million years (My)] and 147Sm–144Nd [T(1/2) = 106 billion years (Gy)]. Both Sm and Nd are refractory lithophile (prefer silicates over metal) elements, whose relative abundances should not be affected by either volatile loss or core formation. The long-lived 147Sm–144Nd system has been widely used to trace planetary-scale processes such as the evolution of the bulk silicate Earth (BSE, defined as all the Earth except for its metallic core) and its chemical differentiation into crust and mantle over Earth’s history. The early epic of Earth’s differentiation is better investigated with the short-lived chronometer 146Sm–142Nd, because of the lack of available samples from Earth’s first 500 My of existence and the sensitivity of most long-lived radioactive systems to resetting by events occurring later in Earth’s history. Recent studies have reported small 142Nd/144Nd excesses in a small number of samples from 3.8 Ga from the Isua Supracrustal Belt, Greenland (1, 2), which is the first evidence for differentiation of the silicate portion of Earth that must have occurred close in time to the well-documented early differentiation of the Moon (3) and Mars (4, 5).

The standard assumption in using this radiometric system to model the geochemical evolution of Earth is that the BSE has Sm/Nd, 142Nd/144Nd, and 142Nd/144Nd ratios approximately equal to those of chondritic meteorites, the building blocks of the planet. Chondritic meteorites show a relatively limited range in Sm/Nd (146Sm/144Nd from 0.1932 to 0.2000), with corresponding 142Nd/144Nd from 0.512525 to 0.512722, when fractionation is corrected to 146Nd/144Nd = 0.7219 (6–8), and the average chondritic values of 0.1966 and 0.512638 (6) have been used to represent the BSE model values for 25 years. Here we present Sm-Nd data from chondrites that overlap previous Sm/Nd and 142Nd/144Nd ratio determinations but indicate a measurable difference of 142Nd/144Nd ratios between chondrites and all terrestrial samples. If the BSE has an Sm/Nd ratio within the range measured for chondrites, the higher-than-chondritic 142Nd/144Nd ratio of terrestrial materials requires that the silicate Earth experienced a global chemical differentiation during the lifetime of 146Sm, resulting in high and complementary low Sm/Nd ratio reservoirs that have remained separate over all of Earth’s history.

Terrestrial Sm-Nd evolution. The most active volcanic system on Earth, the global ocean ridge system, erupts magmas with 142Nd/144Nd ratios considerably higher than any value measured for bulk chondritic meteorites. This value reflects the high Sm/Nd ratio and the general depletion of the oceanic mantle in those elements that selectively partition into melts (the so-called incompatible elements), because Nd is more incompatible than Sm. The incompatible element depletion of the mid-ocean ridge basalt (MORB) source generally is modeled as resulting from the extraction of the incompatible element–rich continental crust over Earth’s history. To explain the high Sm/Nd in the MORB source purely by continental crust extraction from a mantle that initially had chondritic Sm/Nd, only roughly one-third to one-half of the mantle can be as incompatible element–depleted as the MORB source (9–11). The rest of the mantle has typically been assumed to have remained primitive, with a chondritic Sm/Nd ratio. However, some observations are difficult to reconcile with this model (12, 13). For example, recent seismic imaging of the mantle suggests that convective motion may stir at least the upper three-quarters of the mantle (14), if not most of it. Also, the oldest crustal rocks on Earth (dating to 4 Ga) have superchondritic 142Nd/144Nd ratios, requiring an episode of even earlier crustal extraction, although only volumetrically insignificant amounts of this pre–4-Ga crust survive at Earth’s surface today (15, 16). Other evidence that the incompatible element depletion that now characterizes the MORB source mantle is an old feature, not related to the formation of the current continental crust, is the presence of excess 129Xe (derived from the decay of 17-My–half-life 127I) and Pu-fission Xe in the MORB, which indicate that the source must have been outgassed within 50 My of Earth’s formation (17, 18).

Fig. 1. 142Nd/144Nd ratios measured for chondrites and eucrites compared to the La Jolla terrestrial Nd standard (ε142Nd). All chondrites and basaltic eucrites have negative ε142Nd values outside the external analytical error of ± 0.07 ε units (2σ) (shaded area). Cumulate eucrites have positive ε142Nd values in agreement with their high Sm/Nd, resulting from igneous processes on their parent body. The error bars correspond to the internal precision (2σε mean). Terrestrial samples (MORBs, kimberlites, and carbonatites of different ages and collected in diverse locations) measured using the same procedure (27) have been added to demonstrate the significant excess of 0.2 ε units in all the terrestrial material (samples and standard) relative to the mean chondritic value. All terrestrial samples were measured several times using the same procedures as were used for the chondrites. The uncertainties reported on the mean are 2σε.

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Chondritic $^{142}$Nd/$^{144}$Nd. Because of the small variations in $^{142}$Nd/$^{144}$Nd, $^{142}$Nd studies generally use a terrestrial Nd standard to express the $^{142}$Nd anomalies in samples by direct comparison between the two measurements: 

$$
(\frac{\text{Nd}}{\text{Nd}_{\text{standard}}} - 1) \times 10^4
$$

on the assumption that the terrestrial standard has chondritic $^{142}$Nd/$^{144}$Nd. This assumption was derived from measurements that showed no resolvable difference between the $^{142}$Nd/$^{144}$Nd of chondrites and that of the laboratory standard. For example, works by Jacobsen and Wasserburg (6, 7), which are the only published measurements of bulk chondrites that include $^{142}$Nd/$^{144}$Nd data, reported an average $^{142}$Nd of –0.04 ± 0.54 (2σ of the population) with a range from –0.5 ± 0.4 to +0.4 ± 0.4. Amelin and Rotenberg (19) reported measurements of individual chondrules and phosphates separated from a variety of chondrites that provide a $^{142}$Sm-$^{144}$Nd isochron and would yield a present-day value of $\varepsilon^{142}$Nd = 0.01 ± 0.83 for a $^{142}$Nd/$^{144}$Nd of 0.1966. The relatively large errors reflect that these data were either obtained on an older generation of mass spectrometer; measured Nd$^\text{3+}$ instead of Nd$^\text{4+}$, thus requiring an additional correction for oxygen isotope composition and further complicating correction for Ce, Pr, and Sm interference; or were run on small samples. Today, a precision of a few parts per million (ppm) on Nd isotope ratios can be obtained with the new generation of thermal ionization mass spectrometer (2, 20).

We measured the isotopic composition of Sm and Nd on a Thermo-Finnigan Triton thermal ionization mass spectrometer using the Nd$^\text{3+}$ ion beam on samples passed repeatedly through ion-exchange chemistry, in order to reduce Ce and Sm interferences to negligible levels. Details of the analytical procedures and all the raw isotopic data measured on these samples are provided in the supporting online material (21). All of the analyzed chondrites and basaltic eucrites have $^{142}$Nd/$^{144}$Nd ratios lower than the La Jolla Nd standard by between –40 and –8 ppm (Table 1). The deviations compared with the terrestrial reference are clearly significant because the external precision (2σ) measured on repeated runs is lower than 7 ppm for both samples and standards (Fig. 1 and Table 1).

Table 1. Sm-Nd isotope and concentration data for chondrite and eucrite samples. Sm and Nd concentrations were determined by isotope dilution on a spiked aliquot taken after dissolution. $^{143}$Nd/$^{144}$Nd ratios are normalized to 0.511860 for the La Jolla Nd standard. $^{142}$Nd/$^{144}$Nd ratios are expressed in epsilon notation relative to La Jolla Nd (27). For Nuevo Laredo, Pasamonte, and one Sharp, Nd isotopic measurements were made on the same aliquot (only one dissolution) but loaded on separate filament runs in different barrels. The analytical uncertainties correspond to internal within-run precision (2σm). The external reproducibility on the $^{142}$Nd/$^{144}$Nd ratio is about 7 ppm, except for Pasamonte. The lower precision reported for one measurement of Sharp and for Abeé is explained by the small Nd signal. Dissolution in steel-jacket Teflon bombs or closed beakers on a hot plate produced no difference in results. The $^{142}$Sm/$^{144}$Sm of 0.1857±0.0002 ratios relative to the mean of the Sm standard (7 samples) measured for different samples signifies that isotopic ratios have not been significantly affected by neutron flux and electric effects. The internal precision on $^{142}$Sm is lower than 0.3 e. units. Bas. Euc., basaltic eucrite; Cum. Euc., cumulate eucrite.

<table>
<thead>
<tr>
<th>Meteorite name</th>
<th>Class</th>
<th>Sm</th>
<th>Nd</th>
<th>$^{147}$Sm/$^{144}$Nd</th>
<th>$^{143}$Nd/$^{144}$Nd ± 2σm</th>
<th>$^{142}$Nd/$^{144}$Nd</th>
<th>$^{144}$Sm/$^{144}$Sm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Allende*</td>
<td>CV3</td>
<td>0.3147</td>
<td>0.9858</td>
<td>0.1930</td>
<td>0.512573 ± 0.000003</td>
<td>–0.26 ± 0.07</td>
<td>–0.47</td>
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<tr>
<td>Allende</td>
<td>CV3</td>
<td>0.3238</td>
<td>1.010</td>
<td>0.1938</td>
<td>0.512563 ± 0.000002</td>
<td>–0.34 ± 0.05</td>
<td>–0.55</td>
</tr>
<tr>
<td>Allende</td>
<td>CV3</td>
<td>0.3128</td>
<td>0.9802</td>
<td>0.1929</td>
<td>0.512558 ± 0.000003</td>
<td>–0.31 ± 0.07</td>
<td>–0.64</td>
</tr>
<tr>
<td>Allende</td>
<td>CV3</td>
<td>0.512560</td>
<td>0.000003</td>
<td>0.19</td>
<td>0.512620 ± 0.000003</td>
<td>–0.08 ± 0.08</td>
<td>0.19</td>
</tr>
<tr>
<td>Bruderheim</td>
<td>L6</td>
<td>0.1746</td>
<td>0.5399</td>
<td>0.1955</td>
<td>0.512620 ± 0.000003</td>
<td>–0.08 ± 0.08</td>
<td>0.19</td>
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<tr>
<td>Dhaajal</td>
<td>H3.8</td>
<td>0.2012</td>
<td>0.6225</td>
<td>0.1954</td>
<td>0.512621 ± 0.000003</td>
<td>–0.19 ± 0.07</td>
<td>–0.58</td>
</tr>
<tr>
<td>Richardton*</td>
<td>H5</td>
<td>0.1912</td>
<td>0.5952</td>
<td>0.1942</td>
<td>0.512622 ± 0.000003</td>
<td>–0.16 ± 0.08</td>
<td>–2.17</td>
</tr>
<tr>
<td>Richardton</td>
<td>H5</td>
<td>0.512632</td>
<td>0.000004</td>
<td>0.19</td>
<td>0.512632 ± 0.000004</td>
<td>–0.20 ± 0.10</td>
<td>–2.60</td>
</tr>
<tr>
<td>Sharp</td>
<td>H3.4</td>
<td>0.1855</td>
<td>0.5722</td>
<td>0.1959</td>
<td>0.512624 ± 0.000006</td>
<td>–0.30 ± 0.15</td>
<td>–1.61</td>
</tr>
<tr>
<td>Sharp</td>
<td>H3.4</td>
<td>0.1901</td>
<td>0.5920</td>
<td>0.1951</td>
<td>0.512622 ± 0.000002</td>
<td>–0.16 ± 0.04</td>
<td>–0.01</td>
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<tr>
<td>Abeé</td>
<td>EH4</td>
<td>0.1460</td>
<td>0.4640</td>
<td>0.1903</td>
<td>0.512586 ± 0.000008</td>
<td>–0.40 ± 0.17</td>
<td>–0.20</td>
</tr>
<tr>
<td>Bééba</td>
<td>Bas. Euc.</td>
<td>1.699</td>
<td>5.284</td>
<td>0.1944</td>
<td>0.512597 ± 0.000002</td>
<td>–0.24 ± 0.05</td>
<td>0.09</td>
</tr>
<tr>
<td>Nuevo Laredo</td>
<td>Bas. Euc.</td>
<td>2.661</td>
<td>8.283</td>
<td>0.1942</td>
<td>0.512602 ± 0.000003</td>
<td>–0.19 ± 0.06</td>
<td>–0.34</td>
</tr>
<tr>
<td>Pasamonte</td>
<td>Bas. Euc.</td>
<td>2.293</td>
<td>7.126</td>
<td>0.1945</td>
<td>0.512598 ± 0.000003</td>
<td>–0.27 ± 0.05</td>
<td>–0.25</td>
</tr>
<tr>
<td>Binda</td>
<td>Cum. Euc.</td>
<td>0.3566</td>
<td>0.9857</td>
<td>0.2187</td>
<td>0.513319 ± 0.000002</td>
<td>0.08 ± 0.05</td>
<td>0.60</td>
</tr>
<tr>
<td>Moana</td>
<td>Cum. Euc.</td>
<td>0.1575</td>
<td>0.3772</td>
<td>0.2523</td>
<td>0.514937 ± 0.000004</td>
<td>0.57 ± 0.09</td>
<td>–0.54</td>
</tr>
<tr>
<td>Moore County</td>
<td>Cum. Euc.</td>
<td>0.7071</td>
<td>2.082</td>
<td>0.2053</td>
<td>0.512996 ± 0.000005</td>
<td>0.05 ± 0.06</td>
<td>–0.54</td>
</tr>
</tbody>
</table>

* Dissolution in steel-jacket Teflon bomb.
to chondrites are not scarce; they are ubiquitous. Two explanations can account for this observation: (i) the BSE actually has an Sm/Nd ratio higher than that measured for chondrites, or (ii) all terrestrial rocks derive from a mantle reservoir with a high Sm/Nd ratio formed during the short lifetime of $^{146}$Sm, that is, in the first 300 My of solar-system history. Given the refractory lithophile nature of both Sm and Nd and the small range of Sm/Nd ratios measured in different types of chondrites, there is no obvious reason to expect the BSE to have a higher-than-chondritic Sm/Nd ratio, but this option is not impossible. Regardless of whether the high $^{142}$Nd/$^{144}$Nd ratios of terrestrial rocks reflect a nonchondritic Sm/Nd ratio in the BSE or an early differentiation event, the result requires major revision of models of the evolution and structure of Earth’s interior.

The evolution of the 20-ppm excess in the $^{142}$Nd/$^{144}$Nd ratios observed in terrestrial rocks requires a superchondritic Sm/Nd ratio, the magnitude of which depends on when the high Sm/Nd ratio was formed. The later the high Sm/Nd reservoir formed, the higher its Sm/Nd ratio must be to evolve the observed excess in $^{142}$Nd/$^{144}$Nd value compared with that of chondrites. Because the Sm/Nd ratio also affects evolution of the $^{142}$Nd/$^{144}$Nd value, any formation time for this reservoir more than 30 My after solar system formation would require an Sm/Nd ratio so high that the reservoir would evolve a $^{142}$Nd/$^{144}$Nd value higher than that measured in MORBs (Fig. 3). For formation times of 4.562 and 4.537 Ga, the $^{147}$Sm/$^{144}$Nd ratios needed to evolve a 20-ppm excess in $^{142}$Nd/$^{144}$Nd value over the average chondritic value in our data are 0.209 and 0.212, respectively, which translate into present-day $^{143}$Nd/$^{144}$Nd ratios of 0.513041 (e$_{143}$Nd = +8.4) and 0.513129 (e$_{143}$Nd = +10.1). These values overlap the range of Nd isotopic compositions measured in MORBs and also pass through the data for Archean samples (Fig. 4A).

**Size and composition of early terrestrial reservoirs.** Because all terrestrial rocks analyzed so far have $^{144}$Nd/$^{142}$Nd values higher than our chondritic values, including the Isua samples, which are higher still, the early-formed high Sm/Nd reservoir inside Earth must have been the primary source for crust formation throughout at least the last 4 Gy of Earth’s history. Constraints on the size and composition of the high Sm/Nd reservoir (termed here the “early depleted reservoir” or EDR), and its low Sm/Nd complement (termed the “early enriched reservoir” or EER) can be derived by considering the effect that the continental crust extraction would have on the composition of the mantle from which it was derived. Models relating depletion of the MORB mantle to extraction of the continental crust (9–11) use the mass balance equation of elements between continental crust and depleted mantle

$$[x]_{CC} \times M_{CC} + [x]_{MORBs} \times M_{MORBs} = [x]_{BSE} \times M_{BSE}$$

where $[x]$ represents the concentration of element $x$ in the continental crust (CC), MORB source mantle (MORBs), and BSE, and $M$ is the mass of these reservoirs. Given estimates for the average composition of continental crust (29), the depleted mantle (30), the BSE (31), and the known mass of the continental crust, this equation can be solved, providing the one-third to one-half value mentioned earlier (9–11). However, the nonchondritic $^{142}$Nd/$^{144}$Nd of all terrestrial rocks requires a new mass balance equation

$$[x]_{CC} \times M_{CC} + [x]_{MORBs} \times M_{MORBs} = [x]_{EDR} \times M_{EDR}$$

The Sm/Nd ratio needed to evolve the superchondritic terrestrial $^{142}$Nd/$^{144}$Nd results in a present-day $^{142}$Nd/$^{144}$Nd value close to the value measured for the MORB. Therefore, the extraction of continental crust from this already-depleted reservoir (the EDR) cannot have greatly increased the Sm/Nd ratio of the MORB source; otherwise its $^{143}$Nd/$^{144}$Nd value would have evolved to higher values than those observed for any terrestrial rock.

In order to minimize the effect of continental extraction on the degree of compositional fractionation of the residual mantle, the volume of mantle from which the continental crust was extracted must be large. For example, if the EDR was formed at 4.53 Ga, the $^{143}$Sm/$^{144}$Nd ratio of 0.213 that is needed to evolve the terrestrial $^{142}$Nd/$^{144}$Nd value is well matched by adding continental crust (29) to mantle with MORB source composition (30), only if the volume of mantle is as large as the whole mantle overlying the D$^\ast$ layer. In other words, roughly 96% of the mantle must be as incompatible element-depleted as the MORB source, if the continents formed from an EDR with an Sm/Nd ratio high enough to explain the terrestrial $^{142}$Nd/$^{144}$Nd value. Sufficient uncertainty exists in the crustal and MORB source compositions to allow a slightly smaller EDR, but the overriding result is that mantle that is compositionally similar to the MORB source must occupy most of the mantle.

The nonchondritic $^{142}$Nd/$^{144}$Nd value of terrestrial rocks diminishes the importance of a chondritic, or BSE, composition source in the interpretation of the chemical and isotope systematics of Earth. The observation that the Nd-Hf isotope correlation displayed by oceanic basalts does not simultaneously pass through chondritic Hf and Nd isotopic composition is a problem that has led to debate (32). But there is no reason why this correlation should pass through chondritic composition if these reservoirs were derived from an EDR instead of the BSE. Similarly, the correlation between Nd and Sr isotopic composition defined by oceanic basalts has been used to estimate the BSE values of Sr isotopic composition, and hence Rb/Sr ratio, on the as-

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**Fig. 2.** (A) $^{143}$Sm/$^{144}$Nd isochron diagram for chondrites (solid circles) and basaltic eucrites (solid squares) measured in this work. Our data and all previous chondrite measurements (open symbols) scatter along a 4.567-Ga isochron (44). Diamonds are from (6, 7) and triangles are from (8). Errors bars correspond to the internal precision (2σ). (B) $^{146}$Sm-$^{142}$Nd evolution diagram plots as e$_{142}$Nd versus $^{147}$Sm/$^{144}$Nd. The symbols are the same as in (A). Shown for reference is a line corresponding to $^{142}$Nd/$^{144}$Nd evolution with the estimated solar system initial $^{146}$Sm/$^{144}$Nd of ~0.008 (45–48).
assumption that Earth should have a chondritic Sm/Nd ratio. This assumption is no longer valid, at least for the portion of the mantle that has participated in crust formation. Thus, the BSE concentration of elements such as Rb and Pb, whose volatility and/or siderophile (metallic phase) nature make it unlikely that they are present in chondritic relative abundances in the BSE, depends strongly on the composition of the complementary enriched reservoir produced during early terrestrial differentiation.

A missing reservoir? The lack of terrestrial samples with subchondritic $^{142}$Nd/$^{144}$Nd values indicates either that the BSE has a superchondritic Sm/Nd ratio or that the EER that is complementary to the early-formed EDR has never substantially participated in surface volcanism; thus, its composition is poorly constrained. An estimate of the composition of the EER can be obtained from a similar mass-balance approach on the assumption that EDR + EER = BSE, approximating the EDR composition by using that estimated for the MORB source mantle (30). This approach depends on the accuracy of the BSE composition, which, for the reasons described above, is now robust only for refractory lithophile elements, and on whether the MORB source is an adequate representation of the composition of the EDR. As shown in Fig. 5, if the EER is small, for example, the size of the D' layer, then it must be quite enriched in incompatible elements to explain the magnitude of incompatible element depletion in the EDR. If the EER is as large as the mantle beneath 1600 km (14), then it will be considerably less enriched than a D' EER, but still enriched by about a factor of 2 in most incompatible elements as compared with the BSE. The EER, regardless of its size, would thus contain about 43% of Earth's U, Th, and K, leading to ∼9 TW of heat production. The lack of a signature of any low $^{142}$Nd/$^{144}$Nd reservoir in crustal rocks suggests that the EER could reside at the base of the mantle and has not participated in producing rocks sampled at Earth's surface. As such, there is no compelling reason to assume that mantle convection is layered at mid-mantle depths based on the Sm-Nd data, but that there is a convective boundary at great depth in the mantle that has effectively isolated the small EER part of the mantle, unless the BSE has a nonchondritic composition. A similar model for a deep enriched reservoir and its implications for high heat production and for the rare gas evolution of Earth has recently been presented (18). In these models, the EER could serve both as a heat source for plume formation (33) and as a warm blanket to keep the outer core molten through Earth history and provide the energy needed to drive the geodynamo (34).

How the EER formed is suggested by the smooth BSE-normalized incompatible-element patterns calculated here for the EER. Most of the elements shown in Fig. 5 are refractory...
lithophile elements, and this observation is unlikely to be affected by revisions in BSE concentrations, except for elements such as Rb and Pb. The order of elements on the x-axis is based on the relative incompatibility of these elements during shallow melting of the mantle (57), and thus depends primarily on the partition coefficients of clinopyroxene, a phase stable only in the upper few hundred kilometers of the mantle. The smooth patterns for the EER thus suggest that this reservoir was not formed by fractionation of some high-pressure phase such as perovskite, particularly given the distribution coefficients of trace elements recently measured for both Ca- and Mg-perovskite (35). Instead, a better analog for the EER may be the KREEP (basalts rich in K, rare earth elements, and P) component of the Moon, which is a highly incompatible element–enriched residual liquid believed to have formed at low pressure by near-complete crystallization of a lunar magma ocean (36). If the terrestrial EER also formed at shallow depth, its lack of contribution to surface volcanism suggests that it now must reside in the deep mantle. The sinking of a dense, Fe- and Ti-rich residual liquid like KREEP through the lunar mantle has been modeled (37). A similar event may have happened to a shallow terrestrial EER, taking it from near Earth’s surface to the base of the mantle. Such a dense layer could be stable at great depth and remain poorly mixed with the rest of the mantle over the age of Earth (33). The small deficit in 142Nd (–10 ppm relative to the terrestrial standards) recently reported in a few flood basalts from the Deccan Province (38) could indicate some mixing of the enriched deep material with depleted overlying mantle.

Implications for Mars and Earth’s Moon. The new chondritic 142Nd/144Nd value also requires a revision of the interpretation of the early evolution of other planetary bodies, in particular the Moon and Mars. Martian rocks (as sampled by the SNC meteorites) span a wide range of 142Nd/144Nd values [from +90 ppm to –20 ppm, relative to the terrestrial standard (4)]. Compared with the terrestrial 142Nd/144Nd values, martian 142Sm–144Nd systems allow the determination of the differentiation of the silicate portion of Mars to have occurred as late as 40 My after solar system formation (5). Using instead the new chondritic value for the 142Nd/144Nd ratio requires that this time interval be shortened. Martian core formation dated with 182Hf–182W and mantle differentiation could have occurred simultaneously at ~12 My after solar system formation (5, 39).

Most lunar samples have 142Nd/144Nd ratios close to terrestrial values, with the exception of some high-Ti basalts, which have values that are 20 ppm higher (3). The new chondrite values mean that all lunar samples, like Earth’s, have superchondritic 142Nd/144Nd ratios. This suggests that the Moon formed from a giant impact into an already-differentiated Earth, sampling the depleted reservoir that also materializes after Earth formation. The remain-

References and Notes
17. Materials and methods are available as supporting material on Science Online.
47. The new 142Nd/144Nd value is consistent with the superchondritic initial 142Nd/144Nd value measured for the oldest lunar crustal samples (Fig. 4B) (41–43).
The Toll-like receptors (TLRs) play key roles in activating immune responses during infection. The human TLR3 ectodomain structure at 2.1 angstroms reveals a large horseshoe-shaped solenoid assembled from 23 leucine-rich repeats (LRRs). Asparagines conserved in the 24-residue LRR motif contribute extensive hydrogen-bonding networks for solenoid stabilization. TLR3 is largely masked by carbohydrate, but one face is glycosylation-free, which suggests its potential role in ligand binding and oligomerization. Highly conserved surface residues and a TLR3-specific LRR insertion form a homodimer interface in the crystal, whereas two patches of positively charged residues and a second insertion would provide an appropriate binding site for double-stranded RNA.

Innate immunity is based on an ancient and ubiquitous system of cells and molecules that defend the host against infection. This system can recognize virtually all microbes, using a limited repertoire of germ-line–encoded receptors that recognize broadly conserved components of bacterial and fungal cell walls or genetic material, such as double-stranded viral RNA (1, 2).

The Toll-like receptors (TLRs) are among the most important sensors of the innate immune system (3). The ten known human Toll-like receptors recognize pathogen-associated molecules, such as lipoteichoic acid (recognized by TLR2), lipopolysaccharide (TLR4), flagellin (TLR5), and unmethylated CpG DNA motifs (TLR9). Binding of these ligands to TLRs initiates a series of signaling processes that stimulate and orchestrate the innate and adaptive immune responses (4, 5). Human TLRs are implicated in a number of diseases and, hence, constitute potential therapeutic targets (6, 7).

TLRs are integral membrane proteins located either on the cell surface or in intracellular compartments. Their extracellular or ectodomains (ECDs) are responsible for ligand binding and contain 19 to 25 leucine-rich repeat (LRR) motifs that are also found in a number of other proteins with diverse cellular functions (8).

The intracellular domain, known as the Toll/interleukin-1 receptor homology (TIR) domain, recruits adaptor molecules, such as MyD88, TRIF, and TIRAP, to initiate the signaling process (4, 9).

Human TLR3 is activated by double-stranded RNA (dsRNA) associated with viral infection (10), endogenous cellular mRNA (11), and sequence-independent small interfering RNAs (12). TLR3 is distinct from other TLRs in that it is not dependent on MyD88 but rather on TRIF for signaling (13). Other key features of TLR3 signaling include a requirement for phosphorylation of tyrosine residues in the TIR domain (14) and the involvement of phosphatidylinositol-3 kinase (15). In turn, TLR3 activates genes for secreted antiviral cytokines, such as interferon (IFN-β), and those that encode intracellular, viral, stress-inducible proteins (16).

**Overall structure.** The complete ectodomain of human TLR3 without the N-terminal signal sequence (residues 27 to 700) was expressed using a baculovirus system and purified by Ni-nitrilotriacetic acid (NTA) affinity, ion-exchange, and size-exclusion chromatography.

![Fig. 1. Overall architecture of TLR3 ECD in a ribbon representation. The N-terminal cap region is colored blue; the 23 canonical LRRs are in green; and the C-terminal region is in pink. N-linked sugars (N-acetylglucosamines) that are observed in the electron density maps are shown in ball-and-stick representation, attached to their respective Asn residues. The disulfide bond linking LRRs 2 and 3 is drawn in orange, adjacent to the glycosylation site. (A) Side view of TLR3 with the convex face pointing outwards, the concave face inwards, and the heavily glycosylated side face pointing toward the viewer. (B) View rotated 45° from (A) that highlights the continuous β sheet that forms the concave surface. The position of the large insertion in LRR12 that extends toward the glycosylation-free face is marked with an asterisk.](image-url)