A combined Sm–Nd, Rb–Sr, and U–Pb isotopic study of Mg-suite norite 78238: Further evidence for early differentiation of the Moon

J. Edmundson a,*,1, L.E. Borg b, L.E. Nyquist c, Y. Asmerom d

a Institute of Meteoritics, University of New Mexico, Albuquerque, NM 87131, USA
b Lawrence Livermore National Laboratory, Livermore, CA 94550, USA
c Johnson Space Center, Houston, TX 77058, USA
d Department of Earth and Planetary Sciences, University of New Mexico, Albuquerque, NM 87131, USA

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Abstract

Lunar Mg-suite norite 78238 was dated using the Sm–Nd, Rb–Sr, and U–Pb isotopic systems in order to constrain the age of lunar magma ocean solidification and the beginning of Mg-suite magmatism, as well as to provide a direct comparison between the three isotopic systems. The Sm–Nd isotopic system yields a crystallization age for 78238 of 4334 ± 37 Ma and an initial \( \varepsilon^{143}_{\text{Nd}} \) value of \(-0.27 \pm 0.74\). The age-initial \( \varepsilon^{143}_{\text{Nd}} \) (T-I) systematics of a variety of KREEP-rich samples, including 78238 and other Mg-suite rocks, KREEP basalts, and olivine cumulate NWA 773, suggest that lunar differentiation was completed by 4492 ± 61 Ma assuming a Chondritic Uniform Reservoir bulk composition for the Moon. The Rb–Sr isotopic systematics of 78238 were disturbed by post-crystallization processes. Nevertheless, selected data points yield two Rb–Sr isochrons. One is concordant with the Sm–Nd crystallization age, 4366 ± 53 Ma. The other is 4003 ± 95 Ma and is concordant with an Ar–Ar age for 78236. The 207Pb–206Pb age of 4333 ± 59 Ma is concordant with the Sm–Nd age. The U–Pb isotopic systematics of 78238 yield linear arrays equivalent to younger ages than the Pb–Pb system, and may reflect fractionation of U and Pb during sample handling. Despite the disturbed nature of the U–Pb systems, a time-averaged \( \mu \) value of the source can be estimated at 27 ± 30 from the Pb–Pb isotopic systematics. Because KREEP-rich samples are likely to be derived from source regions with the highest U/Pb ratios, the relatively low \( \mu \) value calculated for the 78238 source suggests the bulk Moon does not have an exceedingly high \( \mu \) value.

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1. INTRODUCTION

The lunar magma ocean (LMO) concept asserts that the mantle and crust of the Moon formed during solidification of a molten body by equilibrium and fractional crystallization (e.g., Smith et al., 1970; Wood et al., 1970; Snyder et al., 1992; Shearer and Papike, 1999). Lunar ferroan anorthosites (FANs) are thought to be floatation cumulates formed during the mid- to late-stage of LMO solidification. Following the formation of FANs, extreme fractionation during the last LMO stage concentrated incompatible elements into the KREEP (high K, rare earth element, and P) component, the signature of which is observed in the magnesium suite (Mg-suite), alkali suite, and KREEP basalts. Therefore, the LMO concept predicts that the FANs formed prior to KREEP, and both were followed by the Mg-suite. However, due to the overlapping crystallization ages determined for FANs and Mg-suite samples, as well as the model ages calculated for the formation age of KREEP, the age of LMO solidification is not precisely constrained by the present data set (Fig. 1).

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The apparent temporal overlap between FAN crystallization ages, KREEP model ages, and Mg-suite crystallization ages may stem from the inherent difficulties of dating these materials. For example, FANs are essentially monomineralic anorthosite, and contain only minor amounts of the mafic minerals required to determine a precise mineral isochron age for the sample. Additionally, the isotopic systematics of the FANs may have been disturbed during the post-crystallization impacts which formed the FAN breccias. Isotopic disturbances in FANs are suggested by their positive initial $^{143}\text{Nd}$ values (Carlson and Lugmair, 1988; Alibert, 1994; Alibert et al., 1994; Borg et al., 1999; Norman et al., 2003), which indicate derivation of FANs from a light rare earth element (LREE) depleted source relative to the Chondritic Uniform Reservoir (CHUR). This is inconsistent with the LMO model crystallization sequence from a Moon of bulk CHUR composition, which has mafic minerals crystallizing first and leaving a LREE-enriched melt from which the FANs formed. In this scenario, FANs should have initial $^{143}\text{Nd}$ values that are zero or slightly negative. It is also inconsistent with the calculated chondritic to LREE-enriched parent magma compositions for the FANs (e.g., Korotev et al., 1980). Therefore, if the measured Sm–Nd isotopic systematics of the FANs are not disturbed, they likely do not represent cumulates from a CHUR-like bulk Moon LMO and thus may not yield the correct age of LMO solidification.

It may be possible to date the minimum age of LMO solidification by dating the late-stage material that formed during LMO solidification, KREEP. Unfortunately, KREEP is only a component of some lunar rocks, and thus cannot be dated directly. Model ages for KREEP have been generated using the Sm–Nd and Rb–Sr isotopic systematics of KREEP-rich rocks and breccias, but are highly depen-
dent on the assumptions of the models. For example, an initial Sr isotopic composition for KREEP must be assumed for Rb–Sr model ages (e.g., Nyquist, 1977). It is also necessary to estimate the Rb abundance in KREEP using a combination of assumed Sm/Rb ratios because of the volatility behavior of Rb and its potential loss during impact (Nyquist et al., 1972, 1973, 1974; Nyquist, 1977; Palme, 1977; Nyquist and Shih, 1992). Additionally, the ratio of parent to daughter elements in KREEP calculated using differentiation models are subject to uncertainties in the choice of partition coefficients, and this may inadvertently change the modeled age of KREEP formation. The shaded area of Fig. 1 shows that different assumptions made for different isotopic systems have yielded KREEP model ages that differ by almost 200 Ma (Lugmair and Carlson, 1978; Nyquist and Shih, 1992).

One emerging difficulty with the KREEP model ages, specifically those estimated from the $^{147}\text{Sm}^{143}\text{Nd}$ isotopic system, is the assumption that KREEP is a product of the differentiation of a body with a CHUR bulk composition. This CHUR composition is assumed to be a well-defined, homogeneous starting composition for planetary bodies. However, recent papers (e.g., Boyet and Carlson, 2005; Rankenburg et al., 2006; Nyquist et al., 2008) indicate that specific CHUR values may need to be revised. For example, the howardite–eucrite–diogenite (HED) parent body has an initial $e_{\text{Nd}}$ value of $+0.87 \pm 0.25$ (Nyquist et al., 2004) relative to CHUR, indicating that either the majority of samples originated in reservoirs with super-chondritic $^{147}\text{Sm}^{144}\text{Nd}$ ratios, or the $^{147}\text{Nd}^{144}\text{Nd}$ ratio of CHUR needs to be revised (Nyquist et al., 2008). A greater initial $^{144}\text{Nd}^{144}\text{Nd}$ ratio than CHUR ($e_{\text{Nd}}$ of approximately $+1$) was determined for Precambrian terrestrial samples by Nägler and Kramers (1998), supporting the HED parent body initial $e_{\text{Nd}}$ value. In addition, Rankenburg et al. (2006) determined that lunar samples had chondritic $e_{\text{Nd}}$ values that are 20 parts per million less than terrestrial samples. Caro et al. (2008) claim that there is a difference in $^{143}\text{Nd}^{144}\text{Nd}$ ratios between the Earth, Moon, Mars, and the asteroid belt, which is not due to imprecise measurements in previous studies, but is due to a higher $^{147}\text{Sm}^{144}\text{Nd}$ ratio near the sun during accretion. If this is true, the positive initial $e_{\text{Nd}}$ values of some of the oldest FANs may be explained. However, the geochronology community has not yet adopted a new standard bulk composition for the Sm–Nd isotopic system. Some studies (e.g., Amelin and Rotenberg, 2004) still support the determined CHUR values. Therefore, the CHUR values of $^{143}\text{Nd}^{144}\text{Nd} = 0.512638$ and $^{147}\text{Sm}^{144}\text{Nd} = 0.1967$ will be employed for this study.

Further constraints, independent of bulk composition, can be placed on the minimum solidification age of the LMO by dating the oldest Mg-suite sample. The most reliable system commonly used to date the Mg-suite is $^{143}\text{Sm}^{144}\text{Nd}$ because it seems to be the most resistant to disturbance associated with the late-heavy bombardment. The oldest $^{143}\text{Sm}^{144}\text{Nd}$ date determined on a Mg-suite rock to date is 4460 ± 70 Ma (15445, 17; Shih et al., 1993). However, this sample has a positive initial $e_{\text{Nd}}$ value, which does not agree with the derivation of the sample from a LREE-enriched source relative to CHUR. The positive initial $e_{\text{Nd}}$ value determined for 15445,17 was the impetus for discussing a positive initial $e_{\text{Nd}}$ value of approximately $+1$ for the Moon by Shih et al. (1993). Alternatively, the age and initial $^{143}\text{Nd}^{144}\text{Nd}$ ratio of this sample may indicate a misinterpreted bulk initial $^{147}\text{Nd}^{144}\text{Nd}$ ratio for the Moon (e.g., $e_{\text{Nd}} \sim +1$, Shih et al., 1993; $e_{\text{Nd}} = +0.87$, Nyquist et al., 2006). If it is true that the bulk Moon has an initial $e_{\text{Nd}}$ that is positive, then 15445,17 may be the oldest Mg-suite sample dated. On the other hand, the age and initial of 15445,17 may indicate a LREE-depleted, previously differentiated reservoir parental to the KREEP-rich Mg-suite (e.g., Shih et al., 1993). Due to the multiple conclusions that may be drawn from the 15445,17 data, and the fact that this sample is often assumed to indicate the onset of Mg-suite magmatism, it is important to look at alternative samples that may further indicate the timing of KREEP formation or Mg-suite magmatism (e.g., 78238).

Magnesium-suite norite 78238 comes from the same Apollo 17 Station 8 boulder that has yielded some of the oldest ages for the Mg-suite rocks. The boulder is the parent to samples 78235, 78236, 78238, 78255, and 78256, and represents a single crystallization event (e.g., Jackson et al., 1975). Fig. 1 shows that ages previously obtained for samples from the Station 8 boulder using different isotopic systems range from 4110 ± 20 to 4430 ± 50 Ma (Nyquist et al., 1981; Aeschlimann et al., 1982). The multiple ages obtained for rocks from the Station 8 boulder indicate that the isotopic systematics are disturbed and likely reflect a variety of processes including igneous crystallization and impact metamorphism. There are two preferred $^{147}\text{Sm}^{144}\text{Nd}$ ages and initial $e_{\text{Nd}}$ values obtained for 78236 by Carlson and Lugmair (1981) and Nyquist et al. (1981). The difference in the ages and initial $e_{\text{Nd}}$ values reported by the two groups stems from the interpreted isotopic resetting due to shock noted in the individual mineral fractions by Nyquist et al. (1981) and subsequently removed from the calculated isochron. Thus, the ages and initial $e_{\text{Nd}}$ values presented by Carlson and Lugmair (1981) and Nyquist et al. (1981) for 78236 are sufficiently different to warrant further study of the norites from the Station 8 boulder. The goal of this study is to define the age of this episode of Mg-suite norite formation using the Sm–Nd, Rb–Sr, and U–Pb isotopic systems and thereby constrain the age of LMO solidification and the beginning of Mg-suite magmatism. The same mineral fractions are used for all three systems to make each mineral fraction directly comparable in all three systems. In addition to determining the age of the norite, we hope to precisely determine the initial Nd, Sr, and Pb isotopic compositions of 78238 to better understand the primordial fractionation events associated with LMO solidification.

2. PETROLOGY OF THE STATION 8 BOULDER

The Apollo 17 Station 8 boulder was found at the base of the Sculptured Hills on the southwestern edge of the Serenitatis basin (Jackson et al., 1975). The boulder is approximately 0.5 m in diameter, has a subophitic cumulate texture, and is coated in glass. Samples from the boulder
are composed of approximately 50% chromian bronzite (En$_7$fes$_3$Wo$_{3}$) and 50% plagioclase (An$_{91-97}$), with trace phases such as clinopyroxene, high-Si glass, K-feldspar, troilite, Fe metal, oxides, and phosphates (e.g., Jackson et al., 1975; Sclar and Bauer, 1975; Table 1). Crystallization of the Station 8 boulder norites occurred with both orthopyroxene and plagioclase on the liquidus (Jackson et al., 1975). Jackson et al. (1975) postulated that the glass rind covering the boulder likely formed during ballistic transport. Winzer et al. (1975) noted that the composition of the glass rind matches the whole rock composition of the Station 8 boulder to a first order, which supports the formation of the rind during an excavation (shock metamorphism) event.

The minerals in the norite also show evidence of shock (e.g., Nyquist et al., 1981). Sclar and Bauer (1975) noted that the larger pyroxene grains have undulatory extinction, mosaic, and planar deformation features. Carlson and Lugmair (1981) noted that the pyroxene grains were heavily fractured due to shock, which made the grains break apart easily during crushing as well as during magnetic separation and hand-picking procedures. Additionally, maskelynite (glassy plagioclase) formed from the plagioclase during a metamorphic event in which the boulder experienced shock pressures between 30 and 50 GPa and temperatures above 800 °C (Duke, 1968; Arndt and Gonzalez-Cabeza, 1981; Nyquist et al., 1981).

Compositional effects of shock have been observed in the pyroxene and plagioclase grains within the Apollo 17 Station 8 boulder impact melt (e.g., Sclar and Bauer, 1975). The pyroxenes in the impact melt veins of 78235 have a relatively high Al$_2$O$_3$ content (19–25%), and the plagioclase in the veins is relatively enriched in Fe and Mg when compared to the intact cumulate plagioclase (Sclar and Bauer, 1975). The increase in Fe and Mg in plagioclase within the veins is interpreted to be the incorporation of small amounts of Fe metal and Mg-rich orthopyroxene into the plagioclase during shock (Sclar and Bauer, 1975; El Goresy et al., 1976; Mehta and Goldstein, 1980). Given the major element mobility that can occur in minerals during shock, it is possible that the minor elements, such as those of radiogenic isotopic systems, can be mobilized and redistributed during shock as well. These processes most likely account for the range in ages previously reported for samples from the Station 8 boulder.

### 3. ANALYTICAL METHODS

The chip of norite 78238 used in this study appeared to have an igneous texture, and did not show obvious signs of shock (e.g., shock melt veins) under visual inspection. However, the plagioclase was devitrified into maskelynite. Approximately 1.5 g of 78238 was crushed with a sapphire mortar and pestle. The grains were sieved in a stainless steel container through 100, 200, and 325-size nylon meshes. The 100–200 mesh fraction was then separated magnetically into six different mineral fractions (Fig. 2). The six mineral fractions were then hand-picked under a microscope at 94× magnification to the highest possible purity. Physical descriptions of the mineral fractions analyzed appear in Table 2. The hand-picked fractions were leached with 0.5 N acetic acid, rinsed with 4× quartz distilled water, and then sonicated with 2 N HCl for 10 min. The HCl leachate was removed from the residue fractions and not analyzed. Digestion of the residues was completed with small amounts of ultra-pure concentrated HF and HNO$_3$, and 6 N HCl acids. The mineral fractions were spiked with mixed $^{149}$Sm–$^{150}$Nd, $^{87}$Rb–$^{88}$Sr, and $^{233}$U–$^{236}$U–$^{205}$Pb tracer solutions. The initial chemical separation (U and Pb from Rb + Sr + REE) was completed at the Radiogenic Isotope Laboratory at the University of New Mexico (UNM), using Eichrom analytical grade anion exchange resin 1 × 8, 200–400 mesh, in 50, 100, and 250 μL columns and eluting in 4× quartz distilled HCl, HNO$_3$, and HBr. The Rb + Sr + REE cut was transferred to the Radiogenic Isotope Laboratory at the Johnson Space Center (JSC) for the final chemical separation. The JSC chemical procedures involved separation of Rb, Sr, and REE using 2 and 6 N HCl, and AG50×8 resin. Separation of individual REEs was completed with alpha-hydroxyisobutyric acid (α-HIBA) and NH$_4$-form AG50×8 resin. The Rb and Sr isotope analyses were completed with a Finnigan MAT 261 thermal ionization mass spectrometer (TIMS) at JSC. The U, Pb, Sm, and Nd isotopic ratios were determined at UNM using a VG Sector 54 TIMS. Neodymium isotopic measurements are normalized to $^{146}$Nd/$^{144}$Nd = 0.7219. Neutron fluence effects were measured by analyzing a separate whole rock sample for the Sm isotopic composition. The following blank corrections were incorporated into the data reduction calculations: Rb = 10 pg, Sr = 40 pg, Sm = 20 pg, Nd = 20 pg, and U = 3 pg. The Pb blank corrections varied for each sample, from 12 to 70 pg, and were dependent on the amount of mixed $^{87}$Rb–$^{88}$Sr tracer solution added. Because of the low abundance of $^{204}$Pb in lunar materials, Pb analyses were conducted by measuring $^{204}$Pb in a Daly

### Table 1

<table>
<thead>
<tr>
<th>Samples</th>
<th>78235,6</th>
<th>78235,49</th>
<th>78236,8</th>
<th>78238-d</th>
</tr>
</thead>
<tbody>
<tr>
<td>Orthopyroxene</td>
<td>51</td>
<td>51.2</td>
<td>53.6</td>
<td>32</td>
</tr>
<tr>
<td>Clinopyroxene</td>
<td>0.01</td>
<td>0.01</td>
<td>0.6</td>
<td>&lt;0.1</td>
</tr>
<tr>
<td>Plagioclase</td>
<td>48</td>
<td>48.2</td>
<td>39.2</td>
<td>68</td>
</tr>
<tr>
<td>High-Si glass</td>
<td>0.25</td>
<td>0.25</td>
<td>2.4</td>
<td>Trace</td>
</tr>
<tr>
<td>K-feldspar</td>
<td>Trace</td>
<td>Trace</td>
<td>Trace</td>
<td>Trace</td>
</tr>
<tr>
<td>Phosphate</td>
<td>0.03</td>
<td>0.032</td>
<td>Trace</td>
<td>Trace</td>
</tr>
<tr>
<td>Baddeleyite</td>
<td>Trace</td>
<td>Trace</td>
<td>Trace</td>
<td>Trace</td>
</tr>
<tr>
<td>Zircon</td>
<td>Trace</td>
<td>Trace</td>
<td>Trace</td>
<td>Trace</td>
</tr>
<tr>
<td>Troilite</td>
<td>Trace</td>
<td>Trace</td>
<td>Trace</td>
<td>Trace</td>
</tr>
<tr>
<td>Ilmenorutile</td>
<td>Trace</td>
<td>Trace</td>
<td>Trace</td>
<td>Trace</td>
</tr>
<tr>
<td>Ilmenite</td>
<td>Trace</td>
<td>Trace</td>
<td>Trace</td>
<td>Trace</td>
</tr>
<tr>
<td>Chromite</td>
<td>Trace</td>
<td>Trace</td>
<td>Trace</td>
<td>Trace</td>
</tr>
<tr>
<td>Fe metal</td>
<td>Trace</td>
<td>Trace</td>
<td>Trace</td>
<td>Trace</td>
</tr>
<tr>
<td>Impact melt</td>
<td>4.0</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

a James and Flohr (1983).
b McCallum and Mathez (1975).
c Nyquist et al. (1981).
d Unspecified thin section(s).
multiplier and $^{205}\text{Pb}$ (tracer), $^{206}\text{Pb}$, $^{207}\text{Pb}$, and $^{208}\text{Pb}$ on Faraday collectors. The relative gain between the Daly and Faraday collectors was measured and corrected for in each analysis. Isochrons were calculated with Isoplot 2.49 (Ludwig, 2001) with 2σ errors. The points chosen for use in the isochron calculations are the maximum number for a reasonable linear array. Values for isotopic standards are given in the data tables.

4. RESULTS AND DISCUSSION

4.1. Sm–Nd isotopes

The $^{147}\text{Sm}$–$^{143}\text{Nd}$ isochron obtained for 78238 indicates a crystallization age of 4334 ± 37 Ma with an initial $e^{143}\text{Nd}$ value of $-0.27 \pm 0.74$ (Fig. 3 and Table 3). Despite the presence of impurities such as impact melt in many fractions
Fig. 3. 147Sm–143Nd isochron for 78238. Age of 4334 ± 37 Ma and initial ε143Nd value of −0.27 ± 0.74 determined from all nine data points. Wr = whole rock, Plag = plagioclase, Px = pyroxene, Int = intermediate magnetic separation, GM = glass and mesostasis (interstitial material)-rich fraction, rej = portion rejected during hand-picking, (R) indicates residue after acetic and HCl leaches. Isochron mean standard weighted deviation (MSWD) is 13. Initial ε143Nd calculated using the method of Fletcher and Rosman (1982). Inset indicates deviation of each point from the isochron in squared units.

Table 3

<table>
<thead>
<tr>
<th>Fraction</th>
<th>Wt (mg)</th>
<th>Sm (ppm)</th>
<th>Nd (ppm)</th>
<th>147Sm/144Nd</th>
<th>143Nd/144Nd</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wr 1</td>
<td>12.02</td>
<td>1.419</td>
<td>4.071</td>
<td>0.21074 ± 21</td>
<td>0.513071 ± 20</td>
</tr>
<tr>
<td>Plag-1 (R)</td>
<td>2.98</td>
<td>0.856</td>
<td>4.350</td>
<td>0.11902 ± 48</td>
<td>0.510346 ± 36</td>
</tr>
<tr>
<td>Plag-1 rej (R)</td>
<td>8.09</td>
<td>0.980</td>
<td>4.563</td>
<td>0.12987 ± 17</td>
<td>0.510639 ± 20</td>
</tr>
<tr>
<td>Plag-2 (R)</td>
<td>2.81</td>
<td>0.732</td>
<td>3.136</td>
<td>0.14104 ± 70</td>
<td>0.510953 ± 20</td>
</tr>
<tr>
<td>Mg-Px (R)</td>
<td>15.82</td>
<td>0.936</td>
<td>2.380</td>
<td>0.23782 ± 24</td>
<td>0.513915 ± 20</td>
</tr>
<tr>
<td>Int-Px (R)</td>
<td>21.82</td>
<td>0.603</td>
<td>0.749</td>
<td>0.48644 ± 49</td>
<td>0.520945 ± 20</td>
</tr>
<tr>
<td>Fe-Px (R)</td>
<td>17.17</td>
<td>0.610</td>
<td>0.759</td>
<td>0.48537 ± 63</td>
<td>0.520887 ± 22</td>
</tr>
<tr>
<td>Fe-Px rej (R)</td>
<td>58.78</td>
<td>0.658</td>
<td>1.088</td>
<td>0.36567 ± 37</td>
<td>0.517503 ± 20</td>
</tr>
<tr>
<td>Gm-8 (R)</td>
<td>9.81</td>
<td>1.023</td>
<td>2.553</td>
<td>0.24216 ± 26</td>
<td>0.513975 ± 20</td>
</tr>
</tbody>
</table>

Chemical separations performed at JSC, mass spectrometry performed at UNM. All samples and standards run as NdO. Sm concentration calculated using natural 144Sm/142Sm = 0.560828 (norm), 143Sm/142Sm = 0.515490 ± 2, and 150Sm/142Sm = 0.277439 ± 5 to correct for neutron fluence, ψ = 0.00271 ± 3. Isochrons are calculated using either 2σp (from standard runs) or 2σm (from measured isotopic ratios), whichever is larger.

Error limits apply to last digits and include a minimum uncertainty of 0.5% plus 50% of the blank correction for Sm and Nd added quadratically.

Normalized to 146Nd/144Nd = 0.7219. Uncertainties refer to last digits and are 2σm calculated from the measured isotopic ratios. 2σm = [Σm − μ]²/(n − 1)]1/2 for n ratio measurements m, with mean value μ.

Error limits refer to last digits and are 2σp, 2σp = [Σ(M – μ)²/(N − 1)]1/2 for N measurements M, with mean value π.
plies that the $^{146}$Sm–$^{142}$Nd system does not record the age of the $^{147}$Sm/$^{144}$Nd ratio of 0.178 ± 0.006. All samples in this diagram are consistent with negative $e^{143}$Nd values, light gray polygons = KREEP basalts and olivine cumulate rocks. If the evolved component in both the Moon and Mars represents a late-stage liquid derived by crystallization of a magma ocean, the similar $^{143}$Sm/$^{144}$Nd ratios of the evolved source regions may indicate both lunar and martian magma oceans had similar evolutionary histories.

4.2. Rb–Sr isotopes

The Rb–Sr data for 78238 are disturbed. Nevertheless, two isochrons with apparent age significance can be derived by choosing specific data points (Fig. 5 and Table 4). The oldest age of 4366 ± 53 Ma is defined by the plag-2, whole rock, and Mg-Px fractions. This age is concordant with the Sm–Nd age of 78238, and is also interpreted to represent the crystallization age (C) of the sample. The remaining fractions, as well as the whole rock, define a younger age (M) of 4003 ± 95 Ma. This age is interpreted to represent the time at which the Rb–Sr isotopic systematics of 78238 were disturbed by metamorphism. This conclusion is based on the fact that the youngest Rb–Sr age (M) is concordant with an Ar–Ar age of 4110 ± 20 Ma obtained for a plagioclase separate from 78236 by Aeschlimann et al. (1982).

Since both the Rb–Sr isochrons presented in Fig. 5 contain the whole rock, the disturbance observed in the Rb–Sr isotopic systematics appears to be a rotation of the isochron about the whole rock (e.g., Nyquist et al., 1981, 1991a,b).

Isochron rotation is a likely result of the isotopic re-equilibration of minerals within a rock, without the loss (or gain) of either the parent or daughter element from the system. A mechanism that could explain the rotation of an isochron about the whole rock is post-crystallization isotopic re-homogenization via a metamorphic event. The observation that the plag-2 and Mg-Px fractions define the crystallization age of Fig. 5 suggests that these fractions behave differently during metamorphism than the other mineral fractions. One possible difference between the fractions that lie on the two isochrons which could account for their incongruous responses to metamorphism is the quantity of interstitial material in the fractions. During a shock event, interstitial material can isotopically re-equilibrate within a relatively short amount of time (e.g., Carlson and Lugmair, 1981). This is presumably a result of the
The points used to define the (C) isochron age in the age of metamorphism of 78238 (M). Inset indicates the deviation of quadratically.

with an initial $^{87}\text{Sr}/^{86}\text{Sr}$ of 0.69961 ± 63 Ma and a MSWD of 65. This is interpreted to represent the crystallization age of the sample, and is referred to as (C). The remaining fractions (white dots), as well as the Wr-1, define an age of 4003 ± 95 Ma, age of the sample, and is referred to as (C). The remaining fractions (4343 ± 48 Ma). It is interpreted to represent the crystallization age for the Station 8 boulder determined by this method is 4343 ± 48 Ma, which is concordant with the 4334 ± 37 Ma Sm–Nd isochron age presented above. The fact that the whole rock crystallization age for the Station 8 boulder norites is concordant with the Sm–Nd internal isochron age of 78238 indicates that the Rb–Sr isotopic systematics of the boulder have remained closed during the disturbance that appears to have re-homogenized the Rb–Sr system between various igneous phases.

The initial $^{87}\text{Sr}/^{86}\text{Sr}$ isotopic composition for 78238 from the (C) isochron ($T = 4366 ± 53$ Ma) is 0.699072 ± 0.000044, which is slightly higher than that of Angra dos Reis (ADOR, 0.698970 ± 0.000018; Nyquist et al., 1994) and the Basaltic Achondrite Best Initial (BABI, 0.69898 ± 0.00003; Papanastassiou and Wasserburg, 1969), and is within uncertainty of LUNI (0.69903, Nyquist et al., 1994) and the Basaltic Achondrite Best Initial (BABI, 0.69898 ± 0.00003; Papanastassiou and Wasserburg, 1969), and is within uncertainty of LUNI (0.69903, Nyquist et al., 1994), indicating relatively little production of $^{87}\text{Sr}$ in the source region of 78238. The initial $^{87}\text{Sr}/^{86}\text{Sr}$ for 78238

![Image](86x556 to 283x703)

**Table 4**

Rb–Sr analytical results for 78238.

<table>
<thead>
<tr>
<th>Fraction</th>
<th>Wt (mg)</th>
<th>Rb (ppm)</th>
<th>Sr (ppm)</th>
<th>$^{87}\text{Rb}/^{86}\text{Sr}$</th>
<th>$^{87}\text{Sr}/^{86}\text{Sr}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wr 1</td>
<td>12.02</td>
<td>1.00</td>
<td>30.97</td>
<td>0.09426 ± 47</td>
<td>0.705041 ± 10</td>
</tr>
<tr>
<td>Plag-1 (R)</td>
<td>2.98</td>
<td>1.36</td>
<td>199.3</td>
<td>0.01974 ± 10</td>
<td>0.700434 ± 10</td>
</tr>
<tr>
<td>Plag-1 rej (R)</td>
<td>8.09</td>
<td>2.16</td>
<td>193.2</td>
<td>0.03227 ± 16</td>
<td>0.701065 ± 13</td>
</tr>
<tr>
<td>Plag-2 (R)</td>
<td>2.81</td>
<td>1.51</td>
<td>193.6</td>
<td>0.02250 ± 11</td>
<td>0.700492 ± 10</td>
</tr>
<tr>
<td>Mg-Px (R)</td>
<td>15.82</td>
<td>1.22</td>
<td>31.71</td>
<td>0.11139 ± 56</td>
<td>0.706089 ± 11</td>
</tr>
<tr>
<td>Int-Px (R)</td>
<td>21.82</td>
<td>0.082</td>
<td>0.290</td>
<td>0.8131 ± 41</td>
<td>0.746736 ± 27</td>
</tr>
<tr>
<td>Fe-Px (R)</td>
<td>17.17</td>
<td>0.063</td>
<td>0.256</td>
<td>0.7164 ± 46</td>
<td>0.741230 ± 31</td>
</tr>
<tr>
<td>Fe-Px rej (R)</td>
<td>58.78</td>
<td>0.453</td>
<td>8.91</td>
<td>0.14701 ± 74</td>
<td>0.708622 ± 10</td>
</tr>
<tr>
<td>GM (R)</td>
<td>9.81</td>
<td>3.12</td>
<td>18.23</td>
<td>0.49573 ± 48</td>
<td>0.727904 ± 12</td>
</tr>
</tbody>
</table>

NBS-987 ($N = 6$) 0.710247 ± 10

Chemical separations and mass spectrometry performed at JSC. $^{87}\text{Rb}/^{86}\text{Rb}$ ratio of two runs of the NBS-727 Rb standard was 2.5976 ± 31 (2σ). $^{87}\text{Rb}$ = 1.402 ± 10$^{-11}$ y$^{-1}$.

Isotopic ratios are calculated using either 2σm (from measured isotopic ratios), whichever is larger.

* Error limits apply to last digits and include a minimum uncertainty of 0.5% plus 50% of the blank correction for Rb and Sr added quadratically.

b Normalized to $^{87}\text{Sr}/^{86}\text{Sr}$ = 0.1194. Uncertainties refer to last digits and $2\sigma_m$ calculated from the measured isotopic ratios. $2\sigma_m = (\Sigma m_i - \mu)^2/(n - 1)$

c Uncertainties refer to last digits and $2\sigma_n = (\Sigma (M_i - n\mu)^2)/(N - 1)$

for N ratio measurements $m_i$ with mean value $\mu$.
the incorporation of a component with an enriched age of the Solar System. Estimates of the source region and the source regions formed fairly close to the formation and Tatsumoto, 1991; Premo et al., 1999). This component and Tatsumoto, 1977; Hanan and Tilton, 1987; Premo et al., 1999). Incorporation of a component with a high $^{207}\text{Pb}/^{206}\text{Pb}$ ratio, such as lunar soil Pb, could occur during the shock event(s) that melted the plagioclase in 78238 (e.g., Misawa et al., 2000). However, Papike et al. (1982) noted that the soil at the Apollo 17 Station 8 site had high Sm and Nd abundances, which could also influence the linearity of the 78238 Sm–Nd isochron if the soil were only incorporated into the plagioclase fractions in large quantity. Because the 78238 Sm–Nd isochron does not show evidence for soil addition into the plagioclase fractions it is likely that insufficient soil was added to influence the Sm–Nd isotopic composition of the plagioclase fractions (e.g., the Nd/Pb of the soil is too low), the soil added had a nearly identical Sm–Nd isotopic signature as the plagioclase fractions of 78238, or the high $^{207}\text{Pb}/^{206}\text{Pb}$ component added was elemental Pb mobilized by impact volatility (e.g., Misawa et al., 2000). There is no apparent disturbance to the Pb isotopic systematics of the mafic silicate and whole rock fractions. Accordingly, the line running through the mafic silicate and whole rock fractions in Fig. 7a intersects the ordinate at a $^{207}\text{Pb}/^{206}\text{Pb}$ ratio indicative of an age of 4288 ± 27 Ma, which is concordant with the Sm–Nd crystallization age determined for 78238.

Unfortunately, the $^{235}\text{U}/^{207}\text{Pb}$ and $^{238}\text{U}/^{206}\text{Pb}$ linear arrays obtained for 78238 (not shown) are not concordant with the Sm–Nd, Rb–Sr (C), or the Pb–Pb age. The $^{235}\text{U}/^{207}\text{Pb}$ linear array corresponds to an age for 78238 of 4010 ± 60 Ma and is concordant with the disturbed Rb–Sr (M) age of 4003 ± 95 Ma. The $^{238}\text{U}/^{206}\text{Pb}$ linear array corresponds to an age of 3520 ± 290 Ma and is not concordant with any age determined for this rock. Discordant U–Pb “ages”, specifically those where the $^{238}\text{U}/^{206}\text{Pb}$ age is younger than the $^{235}\text{U}/^{207}\text{Pb}$ age have been reported for lunar troctolite 76535 (Premo and Tatsumoto, 1992), as well as mare basalts 10017 (Gaffney et al., 2007) and 15085 (Unruh and Tatsumoto, 1977). The contribution of terrestrial Pb (chemistry blank) to the fractions may contribute to the discordance between the Pb–Pb and both U–Pb ages due to the large difference between lunar and terrestrial Pb isotopic compositions. However, increasing the blank contribution to the U–Pb isotopic systems will decrease both U–Pb ages equally, resulting in young but discordant $^{235}\text{U}/^{207}\text{Pb}$ and $^{238}\text{U}/^{206}\text{Pb}$ ages. In addition, the leaching procedure followed in this study is identical to that of Gaffney et al. (2007), which shows that all possible surficial terrestrial Pb, as well as some lunar Pb, is removed by this leaching procedure. Therefore, terrestrial contamination is probably not responsible for the separation in $^{235}\text{U}/^{207}\text{Pb}$ and $^{238}\text{U}/^{206}\text{Pb}$ ages observed in 78238. Note that the estimated isotopic composition of terrestrial Pb (Stacey and Kramers, 1975) lies within uncertainty of the 78238 Pb–Pb isochron (Fig. 6). Therefore, mixing the mineral fractions with terrestrial Pb would only draw the points down the Pb–Pb isochron toward the origin. In fact, increasing

![Fig. 6. Pb–Pb isochron for 78238 defining an age of 4333 ± 59 Ma. This age is concordant with the Sm–Nd age and the Rb–Sr (C) age obtained for 78238 during this study. Note that the plagioclase fractions fall off the isochron in the direction of lunar soil compositions (shaded) of Premo et al. (1999). The MSWD of the isochron is 15.948. Nevertheless, the large spread in the data results in a line with the appropriate slope.](Image)
Table 5
U–Pb analytical results for 78238.

<table>
<thead>
<tr>
<th>Fraction</th>
<th>Wt (mg) ID, IC</th>
<th>Pb (ppm)</th>
<th>U (ppm)</th>
<th>$^{206}\text{Pb}/^{238}\text{U}$</th>
<th>$^{207}\text{Pb}/^{235}\text{U}$</th>
<th>$^{208}\text{Pb}/^{235}\text{U}$</th>
<th>Pb blank (pg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wr 1</td>
<td>12.02, —</td>
<td>0.169</td>
<td>0.104</td>
<td>0.004271 ± 2</td>
<td>0.53531 ± 3</td>
<td>275.804 ± 4.768</td>
<td>69</td>
</tr>
<tr>
<td>Plag-1 (R)</td>
<td>2.98, —</td>
<td>0.042</td>
<td>0.021</td>
<td>0.015444 ± 10</td>
<td>1.01504 ± 31</td>
<td>85.307 ± 14.425</td>
<td>38</td>
</tr>
<tr>
<td>Plag-1 rej (R)</td>
<td>8.09, —</td>
<td>0.183</td>
<td>0.195</td>
<td>0.005205 ± 2</td>
<td>0.60272 ± 4</td>
<td>404.704 ± 7.736</td>
<td>56</td>
</tr>
<tr>
<td>Plag-2 (R)</td>
<td>2.81, 22.48</td>
<td>0.081</td>
<td>0.037</td>
<td>0.016830 ± 25</td>
<td>0.82417 ± 68</td>
<td>76.637 ± 6.585</td>
<td>38</td>
</tr>
<tr>
<td>Mg-Px (R)</td>
<td>1.96, 13.86</td>
<td>0.204</td>
<td>0.131</td>
<td>0.002299 ± 3</td>
<td>0.57392 ± 12</td>
<td>579.299 ± 18.762</td>
<td>24</td>
</tr>
<tr>
<td>Int-Px (R)</td>
<td>2.67, 19.15</td>
<td>0.035</td>
<td>0.024</td>
<td>0.007600 ± 19</td>
<td>0.57592 ± 85</td>
<td>187.764 ± 18.418</td>
<td>16</td>
</tr>
<tr>
<td>Fe-Px (R)</td>
<td>2.11, 15.06</td>
<td>0.036</td>
<td>0.019</td>
<td>0.007956 ± 8</td>
<td>0.53769 ± 12</td>
<td>145.062 ± 17.119</td>
<td>14</td>
</tr>
<tr>
<td>Fe-Px rej (R)</td>
<td>7.21, 51.31</td>
<td>0.085</td>
<td>0.061</td>
<td>0.002110 ± 2</td>
<td>0.53599 ± 9</td>
<td>641.411 ± 17.119</td>
<td>14</td>
</tr>
<tr>
<td>Gm (R)</td>
<td>1.18, 8.62</td>
<td>1.120</td>
<td>0.783</td>
<td>0.036484 ± 3</td>
<td>0.53684 ± 10</td>
<td>337.830 ± 4.712</td>
<td>33</td>
</tr>
</tbody>
</table>

Chemical separations and mass spectrometry performed at UNM. Pb ID used for concentration, Pb IC used for Pb isotopic ratios. Pb blank listed is the highest calculated blank for each fraction.

Isochrons are calculated using either $2\sigma_p$ (from standard runs) or $2\sigma_m$ (from measured isotopic ratios), whichever is larger.

$\alpha$ Uncertainties are $2\sigma_m$ calculated from the measured isotopic ratios. $2\sigma_p = (\Sigma m - \mu)^1/(N - 1)^1/2$ for $n$ ratio measurements $m_i$ with mean value $\mu$. Error reported applies to the last significant figure in the ratio.

$\beta$ Error limits include a minimum uncertainty of 0.5% plus 50% of the blank correction for U and Pb added quadratically. Error stated is absolute error.

$\gamma$ Uncertainties are $2\sigma_m$. $\sigma_p = [\Sigma (M_i - \bar{M})^2/(N - 1)]^{1/2}$ for $N$ measurements $M_i$ with mean value $\bar{M}$.

The blank contribution in our data reduction calculations indicates that the Pb–Pb age does not vary with blank contribution. Rather, the error increases with increasing blank contribution.

Other mechanisms that could disturb the U–Pb systematics of 78238 and produce the spread in observed ages are: (1) a change in the $^{238}\text{U}/^{235}\text{U}$ ratio due to neutron capture (Reedy, 1981), (2) impact volatilization and deposition of volatilized lunar Pb, (3) loss of U-series intermediate daughter products (Frisen and Heymann, 1972; Unruh and Tatsumoto, 1977), and (4) preferential loss of Pb relative to U during laboratory handling (e.g., Premo and Tatsumoto, 1992). Neutron capture occurring on the lunar surface has been postulated to change the $^{238}\text{U}/^{235}\text{U}$ ratio from its nominal value of 137.88 (Reedy, 1981). However, the $^{238}\text{U}/^{235}\text{U}$ ratio of 78238 must be lowered to 84.10 in order for the two U–Pb ages to be concordant. Such an extreme deviation from the nominal value, coupled with the measurements of $^{238}\text{U}/^{235}\text{U}$ in lunar materials by Chen and Wasserburg (1981) that indicates no notable deviation from 137.88, does not support a change in the $^{238}\text{U}/^{235}\text{U}$ ratio as the mechanism that caused the difference in U–Pb ages for 78238. Impact volatilization of Pb will increase the $\mu$ values of the fractions if Pb is lost, generating a spread in ages with Pb–Pb $>$ $^{238}\text{U}/^{235}\text{Pb}$ $>$ $^{238}\text{U}/^{206}\text{Pb}$. However, this mechanism decreases all three of the ages relative to the Sm–Nd crystallization age, and is unlikely to be responsible for the disturbed U–Pb systematics in 78238 because the Pb-Pb age is concordant with the Sm–Nd crystallization age. Loss of U-series daughter products, specifically isotopes of Rn gas, can explain the difference in the two U–Pb ages. The longer-lived $^{222}\text{Rn}$ (part of the $^{238}\text{U}$ decay chain, $t_{1/2} = 3.82$ days) may escape more readily from the system than its counterpart on the $^{235}\text{U}$ decay chain ($^{222}\text{Rn}$, $t_{1/2} = 3.96$ s). However, to explain the difference between the crystallization age and the $^{238}\text{U}/^{206}\text{Pb}$ age, preferential diffusion of $^{222}\text{Rn}$ must have occurred over an unreasonably long period (~800 Ma).

Preferential loss of Pb over U during sample handling (e.g., crushing, mineral separating, or leaching) could cause the differences in ages without changing the Pb isotopic composition of the mineral fractions (e.g., Premo and Tatsumoto, 1992). Leachates of 78238 mineral fractions were not analyzed during this study because they are assumed to be mixtures of terrestrial contamination from sample handling, adsorbed lunar surface Pb, and sample Pb (e.g., Premo and Tatsumoto, 1991). However, a 25–28% preferential removal of (lunar) Pb from the mineral fractions could account for the discrepancies in ages between the U–Pb isotopic system and the Sm–Nd, Rb–Sr, and Pb–Pb isotopic systems. Importantly, this process will not influence the Pb–Pb age because Pb isotopes are not fractionated. Thus, it is likely that U–Pb fractionation during sample handling is the disturbance mechanism. It is interesting to note that apparent fractionation of U and Pb during leaching are observed in the most highly shocked lunar and martian samples (e.g., Premo and Tatsumoto, 1992; Misawa et al., 1997). This suggests that leaching-induced fractionation is facilitated by the shock process.

The $\mu$ value for the 78238 source is calculated using a time-averaged, single-stage growth model (4558–4334 Ma) on Fig. 7b, and the mineral fractions that do not appear to be disturbed. The data intersect a single-stage growth curve with a $\mu$ value of 27 ± 30 at 4334 Ma. The relatively large error associated with the $\mu$ value of the 78238 source is the result of the length the 78238 data must be projected to intersect the growth curve. A similar projection on the $^{206}\text{Pb}/^{204}\text{Pb}$ vs. $^{207}\text{Pb}/^{204}\text{Pb}$ diagram yields results that are consistent, but suffers due to both the length of the projection to the growth curve and the large uncertainty on both axes of the diagram. As a result, we prefer the calculation based on the $^{204}\text{Pb}/^{206}\text{Pb}$ and $^{208}\text{Pb}/^{206}\text{Pb}$ ratios in Fig. 7b.

Although the source $\mu$ value estimated for 78238 is significantly lower than the $\mu$ estimated for the single-stage growth of the source of 78235 and 78236 ($\mu$ ~ 508) by Premo and Tatsumoto (1991), it is close to the source $\mu$ for...
mare basalt Yamato-793168 (22 ± 4; Torigoye-Kita et al., 1995) and within error of the source μ of mare basalt Asuka-881757 (10 ± 3; Misawa et al., 1993). It is also within error of the μ value calculated for the time-integrated source region for mare basalt 10017 (70 ± 30; Gaffney et al., 2007), and very close to the μ value calculated for 60025 (35–38) by Premo and Tatsumoto (1992). The relatively low μ value calculated for 78238 contradicts the excessively high μ values for the bulk Moon calculated in previous studies.

High estimates for the μ value of the bulk Moon stem from the high source μ values calculated for KREEP-rich rocks. These estimates tend to be made using the intersection of a best-fit line and a concordia diagram (e.g., Premo and Tatsumoto, 1991). However, the concordia diagram is susceptible to disturbance associated with U and Pb fractionation during sample handling. Disturbance of the U–Pb isotopic systematics by the lunar cataclysm may also result in discordia that intercept the concordia line at shallow angles and therefore result in large uncertainties. We therefore feel that source μ values are most accurately estimated from either Pb–Pb or U–Pb isochron plots that yield ages concordant with crystallization ages determined by other means.

High μ values for KREEP are expected because KREEP is thought to be the latest-stage, most incompatible element-enriched material on the Moon. Any fractionation of U from Pb during LMO solidification by silicates or sulfides is expected to increase the μ value of KREEP (e.g., Neal and Ely, 2002; Gaffney et al., 2007). Therefore, KREEP-rich samples potentially define the maximum bulk Moon μ value. Thus, the μ value estimated for 78238 can be used to constrain the bulk Moon μ value. For example, if there was no fractionation of U and Pb during differentiation of the Moon, then 27 ± 30 is a reasonable estimate for the bulk lunar μ value. If silicates fractionated U from Pb during the formation of KREEP, with U being slightly more incompatible than Pb, then the bulk lunar μ value is less than the maximum 78238 source μ value of 57. If sulfides formed before KREEP during LMO solidification (e.g., Neal and Ely, 2002), removing the majority of Pb from the system and significantly increasing the μ value of KREEP, then the bulk lunar μ value is likely to be significantly less than 57. Regardless of the amount of fractionation of U and Pb that occurred during LMO solidification, the bulk lunar μ value is probably less than 57. Thus, lunar μ values are similar to those estimated for the Earth (<10; e.g., Stacey and Kramers, 1975) and Mars (∼2–5; Chen and Wasserburg, 1986; Borg et al., 2005a; Gaffney et al., 2006).

**5. IMPLICATIONS AND CONCLUSIONS**

The Sm–Nd isochron for 78238 defines an age of 4334 ± 37 Ma with an initial εNd value of 0.27 ± 0.74. The Sm–Nd isotopic systematics of 78238 are consistent with the modeled crystallization sequence of the LMO, and indicates that the Mg-suite is younger than the time of KREEP formation. Sample 78238 therefore places a minimum constraint on the time of LMO solidification. Additional constraints can be placed on the formation age of KREEP by considering the time and initial εNd values of Mg-suite and KREEP basalts. Applying these constraints yields a calculated differentiation age of KREEP of 4492 ± 61 Ma. This model age indicates a relatively early differentiation of a CHUR-like bulk Moon. It further suggests that Mg-suite magmatism started later than previously thought.

The two ages obtained by the Rb–Sr isotopic system for 78238 are 4366 ± 53 Ma (C) and 4003 ± 95 Ma (M). The (C) isochron is concordant with the Sm–Nd age for 78238, and is indicative of crystallization. The (M) isochron is concordant with an Ar–Ar age on a plagioclase separate from 78236. The two different isochrons may be a result of isotopic re-equilibration of interstitial material in the majority of the mineral fractions during a shock metamorphism event.

Multiple “ages” were obtained for 78238 during analysis of the U–Pb isotopic systematics. The Pb–Pb isochron indicated a crystallization age for 78238 of 4333 ± 59 Ma and is concordant with both the Sm-Nd and Rb–Sr (C) ages.
Unfortunately, the U–Pb isotopic systems appear to be disturbed. The $^{238}\text{U}-^{206}\text{Pb}$ linear array indicates an age for 78238 of $4010 \pm 60$ Ma, and the $^{238}\text{U}-^{206}\text{Pb}$ “age” is $3520 \pm 290$ Ma. We hypothesize that fractionation of U and Pb occurred during sample handling and/or leaching of the mineral fractions because other potential disturbances fail to account for all of the observed U–Pb isotopic systematics. A single-stage source $\mu$ value of $27 \pm 30$ was calculated for 78238 and is consistent with a bulk lunar $\mu$ value similar to the Earth and Mars.

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