Timing of crystallisation of the Lunar Magma Ocean constrained by the oldest zircon

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The presently favoured concept for the early evolution of the Moon involves consolidation of debris from a giant impact of a Mars sized body with Earth forming a primitive Moon with a thick global layer of melt referred to as the Lunar Magma Ocean¹. It is widely accepted that many significant features observed on the Moon today are the result of crystallisation of this magma ocean. However, controversy exists over the precise timing and duration of the crystallisation process. Resolution of this problem depends on the establishment of precise and robust key crystallisation time points. We report a 4417±6 Myr old zircon in lunar breccia sample 72215,195, which provides a precisely determined younger limit for the solidification of the Lunar Magma Ocean. A model based on these data, together with the age of the Moon forming giant impact, defines an exponential time frame for crystallisation and suggests formation of anorthositic crust after about 80-85% of the magma ocean was solidified. In combination with other zircon ages the 4417 ± 6 Myr age also
suggests that the very small (less than a few per cent) residual portion of the magma ocean continued to solidify during the following 300-500 m.y. Fractional crystallisation of the Lunar Magma Ocean (LMO) involved the early density-driven separation of mafic cumulates and flotation of a plagioclase-rich lunar crust represented by ferroan-anorthosite. Subsequent crystallisation of ilmenite from the remaining portion of the LMO left a residual liquid enriched in highly incompatible elements. This liquid formed the enriched reservoir referred to as urKREEP (from high concentrations of K, REE, and P).

A precise determination of the timing of fractional crystallisation of the LMO has been inhibited by the susceptibility of Sm-Nd and other systems to the partial resetting during the later thermal pulses associated with the meteorite impacts. As a result, the Sm-Nd mineral isochrons constrained for the ferroan-anorthosite samples show wide spread of ages between 4.56±0.07 Byr (Ref. 3) and 4.29±0.06 Byr (Ref. 4). The best estimate for the age of ferroan anorthosites determined as 4.45±0.07 Myr from the combination of mafic minerals in all analysed samples but excluding plagioclase data that are partially disturbed has another inherited problem as it assumes that all samples have been formed at the same time.

Another way that has been used to constrain the timing of the LMO differentiation is via model ages of rocks derived from different reservoirs in the lunar mantle. In particular, a KREEP-rich source is recognised as an essential part of late stage crystallisation of the LMO, and model ages of urKREEP formation have been estimated as ~4.6 Byr by Rb-Sr analysis of lunar soils, ~4.42 Byr from U-Pb systematics of highlands rocks and a basalt sample and ~4.36 Byr from the Sm-Nd model ages of KREEP samples. An average of model age for KREEP was estimated as 4.42±0.07 Byr (1σ uncertainty). Recent W isotope data on metals from low and
high-Ti mare basalts as well as two KREEP-rich samples\textsuperscript{10} suggest that the last
equilibration of the LMO, which is only possible up to a critical point when about
60\% of the melt is solidified, occurred after 4507 Myr (60 m.y. after formation of the
Solar System). This result is in agreement with \textsuperscript{146}Sm-\textsuperscript{142}Nd model age of the LMO\textsuperscript{10},
which is based on the combined \textsuperscript{147}Sm-\textsuperscript{143}Nd and \textsuperscript{146}Sm-\textsuperscript{142}Nd systems in lunar basalts
and implies a $238_{-30}^{+56}$ m.y. (Ref. 11) to $215_{-21}^{+23}$ m.y. (Ref. 12) time interval for lunar
mantle formation. Despite the general agreement between the model ages determined
using different isotope systems their accuracy is limited by the models and the timing
of LMO remains loosely constrained to the first 250 m.y. of lunar history.

Both isotopic resetting and model dependence problems associated with
numerous previous attempts to place limits on the time of LMO crystallisation can be
avoided by using U-Pb system in zircon\textsuperscript{13,14}, which is well known for its stability
under a variety of extreme conditions. Growth of zircon in melts is governed by
zircon saturation, which can only be achieved in a mafic magma initially enriched in
$\text{Zr}$ (Ref. 15). Consequently, the presence of zircon in the lunar samples is linked to the
initial enrichment of the magma in the KREEP component (i.e., urKREEP must form
on the Moon before zircon can appear in any rock type). Therefore, the oldest zircon
defines a younger limit for the time of urKREEP formation.

Here we report the oldest zircon crystal found on the Moon so far, which is
located in the matrix of Apollo 17 clast-rich impact melt breccia 72215, in the thin
section 72215, 195. The 0.5 mm grain lacks well developed crystal faces and contains
several brittle fractures (Fig. 1), and we thus consider it to be a relict fragment of a
larger grain that was incorporated into the host breccia.

Forty one SIMS U-Pb analyses were made on this grain (Tab. 1, Fig. 2a). The
results indicate a complex pattern of isotope resetting that systematically varies with
the microstructural features of the grain (Tab. 1; Fig. 1). These microstructural
features are a combination of primary magmatic characteristics and different degree of
self-irradiation damage highlighted by the variable birefringence and
cathodoluminescence (CL) emission, as well as deformation patterns revealed by
crystallographic orientation analysis of electron backscatter diffraction (EBSD)
patterns. The observed overall decrease in $^{207}\text{Pb}/^{206}\text{Pb}$ ages correlated with an increase
of the local misorientation determined for each SHRIMP spot$^{16}$ (Fig. 2b), indicates
that this differential resetting of U-Pb system occurred as a result of impact-related
plastic deformation, an interpretation that is consistent with trace element variations
recorded in other deformed zircons$^{16,17}$

All 41 U-Pb analyses are distributed along concordia between 4418±8 and
4331±16 Myr (uncertainties are 2$\sigma$) (Fig. 2a). The four oldest analyses, from
undeformed parts of the grain, form a coherent group on a concordia plot (Fig. 2a)
with concordia intercept at 4420±15 Myr and an average $^{207}\text{Pb}/^{206}\text{Pb}$ age of 4417±6
Myr. We interpret this age as the age of zircon crystallisation. The five youngest
analyses form a coherent group in the $^{207}\text{Pb}/^{206}\text{Pb}$ vs. $^{238}\text{U}/^{206}\text{Pb}$ diagram (Fig. 2a),
defining a concordia intercept age of 4334±10 Myr for the common Pb uncorrected
data and average $^{207}\text{Pb}/^{206}\text{Pb}$ age of 4333±7 Myr for the Stacey-Kramers modern Pb
corrected data. These analyses correspond to areas of moderate luminescence, good
EBSD pattern quality and low U and Th concentrations (Fig. 1). Importantly, these
analyses are also from areas where the deformation bands intersect and/or have high
misorientation, suggesting deformation-related Pb-loss. It is evident that the most
deformed areas of the grain have suffered the greatest Pb loss, and we interpret the
concordia intersection age as a reflection of mobility of the U-Pb system in the grain
during an impact, although the resetting can be incomplete. The remaining
intermediate ages are from areas of moderately-strained parts of the grain, and likely
reflect a partial resetting of U-Pb system.

Our results indicate that the urKREEP source formed by $4417\pm 6$ Myr and it
follows that crystallisation of the LMO was almost completed by this time. The zircon
age is almost 100 Ma older than the age calculated from combined $^{142}\text{Nd}-^{143}\text{Nd}$
systematics of lunar basalts and highland rocks\textsuperscript{11,12}. These later estimates, however,
are based on the assumption that the separate mantle reservoirs have been formed at
the same time and had similar initial isotopic compositions of Nd. This may not be the
case, even for KREEP magmas and the source of high-Ti basalts. Both formed last in
the LMO crystallisation sequence and largely define the slope of combined $^{142}\text{Nd}-$
$^{143}\text{Nd}$ isochrones. Nevertheless, the formation of urKREEP source at $4417\pm 6$ Myr is
in agreement with the age of $4456\pm 40$ Myr determined for the ferroan anorthosite
samples\textsuperscript{5}, even though the ages are not completely resolved within the errors.

A combination of the urKREEP minimum formation age of $4417\pm 6$ Myr and
other data reflecting different stages of LMO evolution allows us to model the history
of magma ocean differentiation and crystallisation on the Moon, and two end-
members are presented (Fig. 3). Both models are constrained by the new $4417\pm 6$ Myr
zircon age, defining a minimum age for formation of Lunar urKREEP at a late stage
in the crystallisation of the LMO. Both are also based on the assumption that the
LMO formed as a result of fast accretion following the giant impact\textsuperscript{1} and, therefore,
the age of LMO formation is similar to the age of the Moon. The best current estimate
of the age of the giant impact based on the Hf-W data is $62_{-10}^{+90}$ m.y. after the
formation of the Solar System\textsuperscript{10}. These data place an older limit of LMO formation of
50 m.y. after the first condensation in the Solar Nebula (i.e. 4517 Myr). A simple
model of LMO evolution (Fig.3, solid line) suggests a sequential fractionation of
olivine $\rightarrow$ orthopyroxene $\pm$ olivine $\rightarrow$ olivine + clinopyroxene $\pm$ plagioclase $\rightarrow$
clinopyroxene + plagioclase $\rightarrow$ clinopyroxene + plagioclase + ilmenite assemblages.

However, the assumption of sequential fractionation of mineral phases throughout the whole LMO is probably an oversimplification because it is likely that: (i) a significant temperature difference would exist between the lower and upper parts of the LMO; (ii) the appearance of different minerals on the liquidus is unlikely to be contemporaneous in different parts of the magma ocean; (iii) convection can prevent effective removal of minerals from the liquid; and (iv) the formation of an insulation lid can change cooling regime of the LMO. A more complex models of LMO crystallisation (Fig.3, dashed line) involves rapid initial cooling of the magma ocean as a result of vigorous turbulent convection$^{18}$, which results in solidification of substantial proportion of LMO without significant fractionation. This was followed by fractionation limited to the relatively thin top layer of the LMO due to much slower cooling resulting from a less vigorous convection regime, and possibly formation of a thermally insulating surface lid.

Nevertheless, both models combined with the available chronological data suggest that ilmenite bearing cumulates precipitated after about 90% of LMO crystallisation, leaving a few percent of residual KREEP melt by 4417±6 Myr. These data suggest that the main volume of the LMO solidified within about 100 m.y. The age distribution patterns obtained for numerous zircon grains from Apollo 17 and 14 breccias$^{14}$ suggest that the residual small volume fraction of the LMO liquid could have cooled slowly over the subsequent 400 to 500 m.y., probably sustained by the internal heating related to radioactive decay. These patterns indicate gradual shrinking of a semi-molten KREEP reservoir towards the centre of Procellarum KREEP terrane$^{14}$, and that by about 4.25 Byr the KREEP reservoir solidified under the area
occupied by the Serenitatis basin, but continued to be active closer to the middle of
Procellarum KREEP terrane near the Imbrium basin until about 3.90 Byr ago.

Assuming that the thickness of the KREEP source is approximately constant
throughout the Procellarum terrane, this accounts for an additional reduction in the
residual proportion of KREEP melt of about 50% by 4.25 Byr.

Despite the precise fixation of the timing of the last stage of LMO
crystallisation by our results, the timing of plagioclase appearance in the
crystallisation sequence remains imprecise. Estimates for the appearance of
plagioclase on the liquidus vary from about 60% to 80% of LMO crystallisation,
depending on the assumed bulk Al content of the LMO\textsuperscript{19,20}. Assuming sequential
crystallisation of minerals (Fig. 3, solid line) and using available geochronological
data for the ferroan anorthosite samples, 70% of crystallisation of LMO is necessary
before plagioclase can become a liquidus phase. In the more complex model (Fig. 3,
dashed line), the lunar crust formed after crystallisation of 80-85% of the LMO.

However, both estimates are within the uncertainties associated with the relatively
imprecise estimate of age of the ferroan anorthosites. The large uncertainty of these
age also results in the large range (anywhere between 20 and 100 my) for the possible
duration of plagioclase flotation. As a result, further refinement of the models awaits
more precise determination of the age of Lunar anorthosite formation.

Methods summary
The sample is a polished thin section of breccia 72215 prepared by NASA. The
microstructure of the zircon was characterized by SEM-based cathodoluminescence
imaging and electron backscatter diffraction (EBSD) mapping using the facilities at Curtin University of Technology, Perth, Western Australia. Collection of EBSD data was processed using the procedures optimised for zircon\textsuperscript{21}. Slip systems were resolved from crystallographic orientation data using simple geometric models of low-angle boundaries\textsuperscript{17}.

U-Pb data were obtained using Sensitive High Resolution Ion Microprobe (SHRIMP) at the John de Laeter Centre of Mass Spectrometry, Curtin University of Technology following the standard analytical procedure described elsewhere\textsuperscript{13}. Pb-U ratios were normalised to the 564 Ma Sri-Lankan zircon CZ3 analysed in a separate mount. Common Pb was corrected using modern Stacey and Kramers lead, following the conclusion that substantial proportion of common Pb in the lunar thin sections results from the surface contamination\textsuperscript{14}. Regardless, of the selection of common Pb for the correction, very low proportion of $^{204}$Pb in the thin section 72215,195 makes the calculated ages insensitive to the uncertainty in the common Pb.

References


Acknowledgements

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**Figure captions**

**Figure 1. Microstructure of the zircon grain from lunar breccia 72215,195.** (a) Optical photomicrograph, cross polarised light showing sector zones and faint compositional growth zones (inset i); (b) panchromatic CL image with superimposed mean U-Pb ages for individual SHRIMP analyses; (c) Map showing variations in EBSD pattern quality (band contrast) from poor (black) to good (white); (d) Map derived from EBSD data showing variations in crystallographic orientation relative to the mean reference orientation (red cross).

**Figure 2. U-Pb SHRIMP data for the zircon from the breccia thin section 72215,195.** a, Tera-Wasserburg concordia diagram. Data are not corrected for the initial Pb. Blue ellipses represent the four oldest analyses; red ellipses represent the five youngest analyses; yellow ellipses represent analyses with intermediate U-Pb ages. b, Age vs. 'local misorientation' value determined at each SHRIMP spot from EBSD map data by calculating the mean misorientation between a central point and its nearest neighbours on an 11x11 pixel grid (i.e., 13.2x13.2 μm area). Local misorientation data were normalised to alpha dose to account for the radiation damage. The resultant local misorientation values are interpreted to reflect lattice distortions associated with crystal-plastic deformation.
Figure 3. LMO crystallisation paths based on the available chronological data.

Solid line projected through the points representing 1) initial formation (100% melt – $^{182}$W age$^{10}$), 2) mean time of lunar crust formation (30% melt – $^{143}$Nd age$^{5}$) 3) KREEP formation (5-7% melt – age from this study), 4) time of cessation of magmatic activity in the Serenitatis region (2.5-3.5% melt – age estimate from zircon distribution patterns$^{14}$); dotted line based on 1) and 2) and the assumption of a turbulent convection in the LMO resulting in the fast initial cooling, yellow circle represents predicted formation of the lunar crust compatible with such fast cooling of the LMO.
Figure 2

**Panel a**

Data-point error ellipses are $2\sigma$

Intercept: $4420 \pm 15$ Ma
MSWD = 0.11; Probability = 0.90

$^{207}\text{Pb}/^{206}\text{Pb}$ Age = $4417 \pm 6$ Ma
MSWD = 0.08; Probability = 0.97

**Panel b**

$^{238}\text{U}/^{206}\text{Pb}$

Intercept: $4334 \pm 10$ Ma
MSWD = 0.08; Probability = 0.97

$^{207}\text{Pb}/^{206}\text{Pb}$ Age = $4333 \pm 7$ Ma
MSWD = 0.05; Probability = 0.99

Error bars are $2\sigma$
Figure 3

Melt remaining (%) vs. Age (my)

- Oldest limit for PI appearance
- Youngest limit for OI-Px cumulates
- ILM cumulate
- KREEP

Data-point error crosses are 2σ
SUPPLEMENTARY MATERIALS

Methods

*Cathodoluminescence*

The panchromatic cathodoluminescence (CL) image was collected using a KE Developments CL system attached to a Philips XL30 SEM at the Microstructural Analysis Facility, Curtin University of Technology, Perth, Western Australia. Operating conditions were 12kV accelerating voltage and working distance of 15mm. The detector sensitivity is in the 330-600nm spectral range.

*Electron backscatter diffraction (EBSD)*

Prior to EBSD analysis, the sample was given an additional polish with 0.06µm colloidal silica NaOH (pH 9.8) suspension using a Buehler Vibromet II polisher for 4 hours to remove the surface damage from previous mechanical polishing, and given a thin (~1nm) carbon coat to reduced the effects of charging in the SEM chamber. Quantitative crystallographic orientation data was collected using EBSD via a Nordlys I detector attached to the Phillips XL30 SEM (20kV accelerating voltage, 20mm working distance, 70° tilt) at Curtin University, and processed using Oxford Instruments Channel 5 (SP9) software following the procedures described in detail for zircon\(^2\). Electron backscatter patterns (EBSPs) were collected (60 ms per frame, 4 frames noise reduction) on a user defined grid (464 x 487 pixels, 1.2µm spacing) and indexed using 8 detected bands; Hough resolution of 65, and match units derived from zircon crystal parameters obtained at 1 atm\(^2\) (Mincryst record: Zircon [2])\(^2\) following detailed assessment of these parameters\(^2\). Some domains of the grain yielded poor quality EBSPs and were unable to be indexed. The average “mean angular deviation” for indexed points is 0.72°. Band contrast is a measure of the EBSP pattern quality (i.e., EBSPs with faint Kikuchi bands yield low band contrast values), and values were obtained from the contrast between the 8 detected bands and the background in a Hough transformation of the EBSPs\(^2\).
Slip systems were resolved from EBSD data using a simple geometric approach that relates the geometry of low-angle tilt and twist boundaries and the dislocations responsible for their formation\textsuperscript{21,25-27}. The map trace of the boundary and the crystallographic dispersion axis were used to reconstruct the 3D boundary orientation, and in turn relate the boundary and dispersion axis orientation to dislocation slip plane and slip direction by assuming end-member tilt boundary models.

**Sensitive high-resolution ion microprobe (SHRIMP).**

Isotopic data were collected using the Sensitive High Resolution Ion Microprobe (SHRIMP II) based in the John de Laeter Centre of Mass Spectrometry, Perth, Western Australia. The SHRIMP methodology follows analytical procedure described elsewhere\textsuperscript{13}. The filtered (O\textsubscript{2}\textsuperscript{-}) beam with intensity between 2 and 3 nA was focused on the surface of samples into \~20 μm spot. Secondary ions were passed to the mass spectrometer operating at a mass resolution (M/ΔM) of \~5000. Each analysis was preceded by a 2 minute raster to remove the Au coating. The peak-hopping data collection routine consisted of five scans through the mass stations, with signals measured by an ion counting electron multiplier. Pb-U ratios were calibrated using an empirical correlation between Pb\textsuperscript{+}-U\textsuperscript{+} and UO\textsuperscript{+}-U\textsuperscript{+} ratios, normalised to the 564 Myr Sri-Lankan zircon CZ3 (Ref. 28). The 0.4 to 1.4% error obtained from the multiple analyses of Pb-U ratio on the standard during individual SHRIMP sessions was added in quadrature to the errors observed in the unknowns. The initial data reduction was done using the SQUID add-in for Microsoft Excel\textsuperscript{29}, and Isoplot\textsuperscript{30} was applied for further age calculations.

The initial Pb correction of lunar samples is complicated by the highly radiogenic Pb compositions of many lunar rocks\textsuperscript{31,320}, which suggest a substantial early loss of Pb from the Moon. A systematic change of $^{206}\text{Pb}/^{204}\text{Pb}$ during SHRIMP analyses of lunar zircon was used to suggest surface contamination as a result of smearing of Pb from the surrounding sample over the zircon surface.
during polishing. However, recent study of 14 thin sections representing different breccia samples from the Apollo 14 and 17 landing sites suggests that although most of the common Pb is a surface contamination, its composition is most similar to the terrestrial Pb (Ref. 14). Therefore, U-Pb analyses obtained for the zircon from the thin section 72215,195 were corrected using modern Stacey and Kramers Pb (Ref. 34). Regardless, of the selection of common Pb for the correction, very low proportion of $^{204}$Pb in the thin section 72215,195 makes the calculated ages insensitive to the uncertainty in the common Pb.

**Internal features of zircon from the breccia thin section 72215,195**

The grain contains several domains, evident from differences in birefringence in cross polarized light (Fig. 1a). These domains have significantly different concentrations of U and Th, which has led to a different degree of self-irradiation damage across the grain. The most U- and Th-rich domain, with U and Th concentrations of ~150 and ~100 ppm respectively and highest Th/U of 0.64 to 0.67 (Tab. 1), also shows very low cathodoluminescence (CL) emission and poor electron backscatter diffraction (EBSD) pattern quality (Fig. 1). Several discrete domains that occur along the edge of the grain, are moderately luminescent and have good EBSD pattern quality (Fig. 1b and c), indicating that the lattice is crystalline. These domains are characterized by low U and Th concentration (~30 to 50 ppm and ~10 to 20 ppm) and the lowest Th/U (0.34 to 0.42, with only one analysis at 0.57). The rest of the grain is dominated by two domains with intermediate U and Th content (~100 to 70 ppm and ~70 to 40 ppm), Th/U (0.56 to 0.60), CL intensity and EBSD pattern quality (Fig. 1b and c). One of these domains records fine scale variations in birefringence (Fig. 1a, insert), interpreted to reflect primary (magmatic) growth zoning with associated minor chemical variation.

Crystallographic orientation analysis reveals that the zircon contains several deformation bands that transect primary zoning and predate brittle fractures (Fig. 1d). Two orthogonal sets of straight discrete and gradational low-angle boundaries accommodate ~12° misorientation across the
grain. The deformation bands are parallel to the crystallographic \( a \)-planes \{010\} of the zircon, have misorientation axes parallel to the \( c \)-axis, and are geometrically consistent with formation by dislocation creep associated with \(<100>\{010\}\) slip\textsuperscript{21}. The deformation bands are geometrically similar to dislocation microstructures reported in experimentally shocked zircon\textsuperscript{35}. We interpret these crystal-plastic deformation microstructures to have resulted from a significant impact, either directly from impact shock, or during ductile flow directly following the impact. The deformation bands appear to continue undeflected through the non-indexed, radiation-damaged areas of the grain, which indicates that the orientation variation predates any significant mechanical weakening from radiation damage in the grain, and therefore occurred early in its history. Crosscutting relationships between the deformation bands and the major chemical domains, identified within the grain, also demonstrate that the observed variation in U concentration and Th/U predate deformation and is the primary growth feature of this zircon.

References


Table 1: U-Pb SHRIMP data for the lunar zircon grain from the breccia thin section 72215,195

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\(\text{a all errors are }\%\ 1\text{ sigma}\)

\(\text{b }^{206}\text{Pb* is radiogenic}^{206}\text{Pb}\)

\(\text{c }\%\ \text{discordance}\)