Fractional crystallization plays a critical role in generating the differentiated continental crust on Earth. However, whether efficient crystal-melt separation can occur in viscous felsic magmas remains a long-standing debate because of the difficulty in discriminating between differentiated melts and complementary cumulates. Here, we found large (~1 per mil) potassium isotopic variation in ~54 strongly peraluminous high-silica (silicon dioxide >70 wt %) leucogranites from the Himalayan orogen, with potassium isotopes correlated with trace elemental proxies (e.g., strontium, rubidium/strontium, and europium anomaly) for plagioclase crystallization. Quantitative modeling requires up to ~60 to 90% fractional crystallization to account for the progressively light potassium isotopic composition of the fractionated leucogranites, while plagioclase accumulation results in enrichment of heavy potassium isotopes in cumulate leucogranites. Our findings strongly support fractional crystallization of high-silica magmas and highlight the great potential of potassium isotopes in studying felsic magma differentiation.

**INTRODUCTION**

Magmatic differentiation induced by continuous crystal-melt segregation plays an important role in shifting the composition of Earth’s continental crust from mafic to felsic (1, 2). However, despite the well-constrained dynamics of crystal settling from mafic magmas (3), it remains debated whether—and, if so, how—efficient segregation of crystals occurs in silicic magmas (4–12), which solidify to form the major components (e.g., granites) of the upper continental crust (13). High-silica (>70 weight % (wt %) SiO₂) liquids are characterized by high viscosities and similar densities to the main crystallizing phases such as quartz and feldspar, meaning that fast and large-scale gravity-driven settling and accumulation of crystals are hard to achieve (5, 6, 10). Alternatively, compaction of crystal mush and hindered settling have been argued as probable mechanisms of separation of high-silica melts and complementary felsic cumulate residues (4, 5, 12). However, unlike mafic magmatic systems where numerous examples of cumulate textures have been observed [e.g., (14)], unequivocal petrological evidence for crystal accumulation in felsic rocks is elusive (15). This may be because either felsic cumulates are rare in the accessible parts of the continental crust (16, 17) or the cumulates are not readily distinguishable from the differentiated high-silica melts possibly due to the trap of interstitial melts in crystal mushes, resulting in similar mineral assemblages and bulk chemical compositions between cumulates and fractionated melts (4, 15). Furthermore, the subtle major elemental variations in evolved high-silica igneous rocks make it difficult to discriminate between strongly and weakly fractionated melts based on bulk rock chemistry such as SiO₂ and MgO (18), which are widely used in tracing the differentiation of mafic-intermediate magmas.

Potassium (K), with two stable isotopes (⁴¹K and ³⁹K), is highly concentrated in high-silica magmas. The compatibility of K is opposite in plagioclase and K-feldspar (19), which are two ubiquitous and major phases in crystallizing assemblages of high-silica magmas. Thus, any change in the modes of plagioclase and K-feldspar will affect the geochemical behavior of K during fractional crystallization of high-silica magmas, leading to the nonsystematic variation in K concentration commonly observed in high-silica igneous rocks [e.g., (18, 20, 21)]. Recent studies revealed large K isotope fractionation among minerals in igneous rocks (22–24). In particular, plagioclase is significantly enriched in ⁴¹K compared to other coexisting rock-forming minerals [e.g., K-feldspar, hornblende, and mica; (23, 24)] due to the decrease in K-O bond length with (Na + Ca)/K in feldspars (25, 26). Accordingly, systematic K isotopic variation might be expected in melts that experience different degrees of plagioclase crystallization. To date, only two studies reported limited K isotope fractionation during magmatic differentiation of basalts (27, 28). This is consistent with the low concentrations of K in the major crystallization phases of mafic magmas (e.g., olivine and pyroxene), leaving the K isotopic system in melts undisturbed. By contrast, scarce data for high-silica granites suggest slightly more variable and lighter K isotopic compositions compared to mafic-intermediate rocks, which was not only previously ascribed to source heterogeneity (29) but may also result from the segregation of isotopically heavy feldspars (28).

To explore the potential of K isotopes in studying the magmatic differentiation of felsic magmas, we performed high-precision K isotopic analysis on a suite of well-characterized leucogranites from the Himalayan orogen (fig. S1). These leucogranites exhibit high-silica and strongly peraluminous characteristics of S-type granites (Fig. 1A and fig. S2) (18, 30). Their parental melts formed via anatectic of granulite- to eclogite-facies metasediments with little evidence for either the presence of mafic rocks in their sources or mixing with mantle-derived or more juvenile mafic melts (30, 31). Therefore, these leucogranites provide an excellent opportunity to investigate the magma chamber processes in an exclusive high-silica magmatic system. We found ~1 per mil (%) K isotopic variation in the Himalayan leucogranite samples, which is most likely produced by plagioclase fractionation and accumulation during magmatic differentiation.
Average UCC = 33), with 41K values > −0.2‰ exclusively 0.04 to 0.35 ± 0.03‰; n varying from average crust-like to significantly higher values (−0.51 ± 41K values $\delta^{41}$ crust [−0.44 ± 0.05‰; (29, 72–75); feldspar-rich sediments (i.e., graywacke, sandstone, and siltstone); (29, 75, 76)]. Data for global granitoids from (29) are plotted for comparison in (A) and (B). The purple areas are the SiO2 content (66.6 ± 2.4 wt %), K2O content (2.80 ± 0.46 wt %), and $\delta^{41}$K value (−0.44 ± 0.05‰) of the average upper continental crust (UCC) (13, 29).

**RESULTS**

Fifty-four well-characterized Cenozoic leucogranites from 11 plutons within the Himalayan orogen, covering a broad range of geographic locations and emplacement ages (table S1 and fig. S1), were measured in this study. These leucogranites are primarily composed of light-colored minerals (>90%), e.g., quartz, feldspar, and muscovite, and are subdivided into a biotite-muscovite (termed “two-mica” hereafter) and a muscovite facies based on the mafic mineral assemblage. Two-mica leucogranites typically consist of quartz, plagioclase, K-feldspar, biotite, and muscovite, while tourmaline and garnet are the major mafic minerals in muscovite leucogranites. The $\delta^{41}$K value varies from −0.72 ± 0.09 to 0.35 ± 0.03‰ in all 54 samples (Fig. 1), which is significantly beyond the range of granitoids reported so far [−0.57 to −0.38‰; (23, 29); see Materials and Methods for the definition of $\delta^{41}$K; its uncertainty is given as 2 SD]. Except for three samples with $\delta^{41}$K values lower than the average upper continental crust [−0.44 ± 0.05‰; (29)], two-mica leucogranites have $\delta^{41}$K values varying from average crust-like to significantly higher values (−0.51 ± 0.04 to 0.35 ± 0.03‰; n = 33), with $\delta^{41}$K values > −0.2‰ exclusively present in samples containing ≤3 wt % K2O (Fig. 1B). By contrast, $\delta^{41}$K values of muscovite leucogranites are overall lower than those of two-mica leucogranites and range from average crust-like to notably lower values (−0.72 ± 0.09 to −0.44 ± 0.09‰; n = 18).

**DISCUSSION**

The large K isotopic variation in the Himalayan leucogranites does not correlate with emplacement age or geographic locations of different plutons (fig. S3A). Instead, $\delta^{41}$K value covaries with whole-rock chemistry, in particular, trace elements such as Rb, Sr, Ba, and Eu/Eu* (Fig. 2 and fig. S3, B to D). Possible processes that can modify K isotopic compositions of Himalayan leucogranites include chemical weathering, source heterogeneity [e.g., (32–34)], incongruent partial melting of mica and feldspar during crustal anatexis [e.g., (35–37)], and crystal-melt separation [e.g., (38, 39)]. Although postemplacement weathering processes could potentially lower the $\delta^{41}$K values of granites (40), 50 of 54 samples are fresh and have chemical index of alteration values [CIA = 50 to 56; see table S1 for definition (41)] within the typical range of unweathered granites. Four samples with slightly elevated CIA values (57 to 61) have very high $\delta^{41}$K values (fig. S4), opposite to the low $\delta^{41}$K value expected for chemical weathering. Thus, chemical weathering cannot be the main cause for the large
K isotopic variation in our samples. The wide range of δ\(^{41}\)K values in the Himalayan leucogranites must result from either source heterogeneity or magmatic processes, which will be discussed in detail below.

**Origins of K isotopic variation in the Himalayan leucogranites**

Mica-rich metapelites and feldspar-rich metagreywackes have been regarded as the dominant source lithologies of the Himalayan leucogranites mainly based on the radiogenic (e.g., Sr, Nd, and Hf) and stable (e.g., O and Si) isotopic resemblance to the Himalayan metasediments (31, 32, 34, 42) and comparable peraluminous compositions to experimental melts of these metasediments (36). Anatexis of these two types of sediments may show distinct chemical and isotopic compositions (32–34, 42). Nevertheless, most sediments have δ\(^{41}\)K values similar to the average upper continental crust with none comparable to the highest δ\(^{41}\)K values observed in the two-mica leucogranites (Fig. 1C). Furthermore, clay/mica-rich sediments and feldspar-rich sediments display no obvious difference in the mean δ\(^{41}\)K value. Last, K isotopes in leucogranites do not correlate with radiogenic isotopes (e.g., Sr and Nd) indicative of provenance and lithology of the Himalayan metasedimentary sources (fig. S5) (32, 34). Therefore, the distinct K isotopic signatures between two-mica and muscovite leucogranites cannot be simply explained by source heterogeneity.

Production of leucogranite by anatexis of metasediments is mainly governed by the following reaction: muscovite + plagioclase + quartz ± H\(_2\)O → melt ± K-feldspar ± biotite ± aluminum-silicate, where the proportions of muscovite and plagioclase contributing to the melt strongly depend on the activity of H\(_2\)O (36). Given the large K isotopic difference between plagioclase and mica (23–26), incongruent melting of muscovite or plagioclase may fractionate K isotopes between melt and its source. Here, we quantitatively evaluated the extent of K isotope fractionation during anatexis using the same nonmodal partial melting model as previously used to quantify Sr and Nd isotopes (see Materials and Methods for details) (43). The results yield <0.07‰ deviation in δ\(^{41}\)K value between melts and the source rock even if a 4% K isotopic fractionation between plagioclase and mica is assumed (fig. S6). This is consistent with the fact that muscovite has K abundance tens of times higher than plagioclase in metasediments (36, 44); thus, mica is the dominant contributor of K to the melt during anatexis. Therefore, partial melting processes cannot generate the ~1‰ δ\(^{41}\)K variation in the Himalayan leucogranites either.

The most likely process responsible for the progressive decrease in δ\(^{41}\)K value from two-mica leucogranite to muscovite leucogranite is the continuous separation of plagioclase during magmatic differentiation, which is the only known phase that could be significantly enriched in heavy K isotopes (23–26). This is supported by the positive correlations of δ\(^{41}\)K value with sensitive proxies for plagioclase crystallization, e.g., Sr and Eu/Eu* (Fig. 2). These positive trends anchor the δ\(^{41}\)K values of the primary leucogranite melts to average crust-like values. A few two-mica leucogranites with the highest Sr and Eu/Eu* do not follow the trends and have much higher δ\(^{41}\)K (>−0.2‰) values than the average upper crust and any potential source rocks (Fig. 1). They most likely represent the plagioclase-rich cumulates, as supported by the enrichment in plagioclase over K-feldspar (fig. S7).

Both fractional and equilibrium crystallization models were applied to assess the influence of crystal-melt separation on the δ\(^{41}\)K values of the Himalayan leucogranites (see Table 1 for model parameters and Materials and Methods for model details). Given that Rb and Sr behave differently during fractionation of mica and feldspar, the Rb/Sr ratio was modeled to further constrain the degree of crystallization. Note that Rb/Sr in the primary melts can vary greatly because of potentially different source lithologies or different melting reactions (32, 33, 45). Therefore, 3 two-mica leucogranites having average crust-like K isotopic compositions (δ\(^{41}\)K = −0.39, −0.41, and −0.44‰Crust) and different Rb/Sr ratios (3.65, 1.13, and 0.48) covering the main Rb/Sr range of Himalayan leucogranites were selected as the primary melts in our models. The modal abundance of the crystallizing assemblage throughout the crystallization was determined using the Rhyolite-MELTS software (Fig. 3). Our modeling results match the K isotopic variation in the Himalayan leucogranites. Cumulates produced during both fractional and equilibrium crystallization have similar δ\(^{41}\)K values, both of which are higher than those of the primary melts and proportional to the modal abundance of plagioclase (Fig. 4 and fig. S7). By contrast, fractional crystallization fits data for the isotopically light leucogranites better compared to equilibrium crystallization (Fig. 4). The isotopically lightest leucogranite samples require ~60 to 90% fractional crystallization, which is close to the optimal (~50 to 70%) crystallinity window for efficient separation of high-silica liquids from crystal mushes (46).

The above crystallization model is simplified as a closed-system crystal-melt separation. Fractional crystallization of high-silica magmas is often accompanied by wallrock assimilation, late-stage fluid exsolution, and magmatic-hydrothermal interaction (47–51). Although these processes could also contribute to the K isotopic variation in the Himalayan leucogranites, their contributions are considered minimum. Country rocks of these Himalayan leucogranites are mainly composed of metasediments, and δ\(^{41}\)K values reported for sediments...
vary from crust-like to lighter K isotopic compositions (Fig. 1C). Assimilation by these sediments cannot explain the elevated $\delta^{41}$K values of those two-mica leucogranites. On the other hand, the Himalayan metasediments have low Rb/Sr (fig. S8); assimilation should have lowered the Rb/Sr of the leucogranites, which is opposite to the higher Rb/Sr for the isotopically lighter muscovite leucogranites (Fig. 4). In addition, these metasediments have K$_2$O similar to or lower than the muscovite leucogranites (fig. S8), implying that an unrealistically high amount of assimilants would be required to alter leucogranite $\delta^{41}$K values. Fluid exsolution and fluid-melt interactions can occur at the advanced stage of magmatic differentiation (48, 50, 51). Partitioning of K between exsolved fluid and granitic melt ($D_{K\text{fluid/melt}}$) is very low in aqueous fluids but linearly increases with the chlorinity of the fluid ($m_{Cl}$) [$D_{K\text{fluid/melt}} = 0.2 \times m_{Cl}$ (52)]. Nevertheless, laboratory experiments and theoretical calculations suggest limited equilibrium K isotope fractionation between Cl-rich fluid and K-rich silicates at the crystallization temperatures of Himalayan leucogranites (fig. S9) (25, 53, 54). Therefore, both country rock assimilation and late-stage fluid activity have minor effects on K isotopes. Crystal-melt separation, therefore, exerts the first-order control on the large K isotopic variation in the Himalayan leucogranites.

Fig. 3. Numerical modeling of progressive crystallization of the Himalayan leucogranites. The modal percentages of crystallizing mineral assemblages during fractional (A to C) and equilibrium (F to H) crystallization of three hypothetical parental leucogranites (09FW56, 09FW144, and 17CN21) were calculated using the Rhyolite-MELTS software. The partitioning coefficient of K ($D_{K\text{crystal/melt}}$) and K isotope fractionation factor ($\Delta^{41}K_{\text{crystal-melt}}$) between bulk crystallizing mineral assemblage and melt for the chosen parental leucogranites used in the fractional (D and E) and equilibrium (I and J) crystallization models are also plotted. See main text for model details.
Understanding the processes of felsic magma differentiation is crucial for studying the origin of granitoids (55), the relationship between volcanic and plutonic felsic magmas (56–58), rare-metal (e.g., Sn, W, Nb, Ta, and Be) mineralization associated with high-silica magmatism (30, 59), and catastrophic rhyolitic eruptions (5). The recognition of crystal fractionation and accumulation in felsic magmas serves as a prerequisite for addressing the above issues. However, evaluating the degree of fractional crystallization of high-silica magmas and finding felsic cumulates are very challenging. For example, trace element signatures such as subchondritic Nb/Ta and Zr/Hf and tetrads (rare earth element) patterns have been regarded as important markers of highly evolved felsic magmas (30, 49), but they are not informative in quantitatively estimating the extent of fractional crystallization. Compatible elements, e.g., Sr, Ba, Eu, and Zr, can be used to identify felsic cumulates because they are enriched in cumulates (15, 56, 60). However, their enrichment is not often obvious, and the inefficient extraction of interstitial melts from the crystal mush further complicates the diagnostic trace element signatures between cumulates and primary melts (4, 15, 56). Our study reveals significant K isotope fractionation during both crystal fractionation and cumulation processes in the Himalayan leucogranites. As plagioclase commonly occurs as a major phase during prolonged differentiation of felsic magmas, K isotopes can provide unprecedented insights into the evolution of felsic magmas. Other nontraditional stable isotope systems such as Ca also display detectable fractionation between plagioclase and other Ca-bearing silicates (e.g., hornblende) in felsic rocks and might be used in a similar way to K isotopes to constrain crystal-melt separation in high-silica magmas (61, 62).

**Potassium isotopic analysis**

Potassium isotopic compositions were determined at the Isotope Laboratory of the University of Washington, Seattle, following a previously established protocol (64, 65). A brief description of this method, together with some minor improvements, was summarized in Materials and Methods.
here. Around 5 to 15 mg of whole-rock powders were successively digested in Teflon beakers with HF-HNO₃ mixture, HCl-HNO₃ mixture, concentrated HNO₃, and 0.5 N HNO₃. To achieve purification of K, an aliquot of solution containing 100 to 200 µg of K was passed through a column filled with 2 ml of Bio-Rad AG50W-X8 (200- to 400-mesh) resin and was eluted using 0.5 N HNO₃. This procedure was performed twice to ensure effective purification with a K yield of >99%. The whole procedural blank contains <10 ng of K, which is negligible compared to the amount of K loaded onto the columns.

The isotopic ratios were measured on a Nu Plasma II multicollector inductively coupled plasma mass spectrometer using the standard-sample bracketing method to correct the instrumental mass bias. The K concentrations of both sample and bracketing standard National Institute of Standards and Technology (NIST) standard reference material (SRM) 3141a were diluted to 3 parts per million (ppm) in 3% HNO₃ before measurement. To overcome the strong isobaric interference of 40Ar+H⁺ on 41K, solutions were introduced into the plasma using a CETAC Aridus II desolvating nebulizer system at the radio frequency power of 700 to 750 W. The 39K and 41K signals were measured simultaneously at an interference-free shoulder in pseudo-high-resolution mode. An ice bucket with constant temperature of ~5°C was used to store the waste acid bottle of the Aridus II, which helps to improve the signal stability and sensitivity. The intensity of ~5°C was used to store the waste acid bottle of the Aridus II.

The isotopic composition of the source rock (41K source) is given by

\[ \delta^{41} K = \left( \frac{^{41} K / ^{39} K}_{\text{sample}} / \frac{^{41} K / ^{39} K}_{\text{NIST SRM 3141a}} - 1 \right) \times 1000 \]

Accuracy and reproducibility were assessed on the basis of the analysis of three granite standards (G-2, GS-N, and JG-1) and one rhyolite standard (RGM-1), which were dissolved and processed through column chemistry together with the samples. Their \( \delta^{41} K \) values agree with the published values (table S2). Multiple analyses on these rock standards also yielded consistent results.

**Modeling K isotope fractionation during anatexis of metapelites**

The melting reaction is described as: muscovite + plagioclase + quartz ± xH₂O → melt ± K-feldspar ± biotite ± aluminum-silicate. The K isotopic composition of the source rock (\( \delta^{41} K_{\text{source}} \)) is given by

\[ \delta^{41} K_{\text{source}} = \frac{\sum_{\alpha=1}^{n} (X_{\text{mineral } \alpha}^0 \times [K_2O]_{\text{mineral } \alpha} \times \delta^{41} K_{\text{mineral } \alpha})}{\sum_{\alpha=1}^{n} (X_{\text{mineral } \alpha}^0 \times [K_2O]_{\text{mineral } \alpha})} \]

where \( X_{\text{mineral } \alpha}^0 \), [K₂O]_{mineral α} and \( \delta^{41} K_{\text{mineral } \alpha} \) are the mass proportion of mineral α in the source rock, K₂O content of mineral α, and K₂O value of mineral α, respectively. The model parameters are presented in table S3.

We assume a constant K₂O content and a \( \delta^{41} K \) value for each mineral during melting processes. The K isotopic composition of melt (\( \delta^{41} K_{\text{melt}} \)) can be calculated using the following equation

\[ \delta^{41} K_{\text{melt}} = \frac{\sum_{\alpha=1}^{n} (X_{\text{mineral } \alpha}^L \times [K_2O]_{\text{mineral } \alpha} \times \delta^{41} K_{\text{mineral } \alpha})}{\sum_{\alpha=1}^{n} (X_{\text{mineral } \alpha}^L \times [K_2O]_{\text{mineral } \alpha})} \]

where \( X_{\text{mineral } \alpha}^L \) is the mass proportion of mineral α entering the melt. The mass proportions of reacting phases entering the melt at different degrees of melting are from table 2 in the work of Patiño Douce and Harris (36). The \( \delta^{41} K \) evolution in melts during partial melting of metasediments at 6 and 10 kbar both without H₂O and with 2 to 4 wt % H₂O is reported in fig. S6.

**Equilibrium crystallization model**

The crystals and melt remain in equilibrium throughout crystallization in this model, so K isotopic compositions of differentiated melt (\( \delta^{41} K_{\text{melt}} \)) and complementary cumulate (\( \delta^{41} K_{\text{cumulate}} \)) can be calculated using the isotope mass balance equations

\[ \delta^{41} K_{\text{melt}} = f^{K}_{\text{melt}} \times \delta^{41} K_{\text{melt}} + (1 - f^{K}_{\text{melt}}) \times \delta^{41} K_{\text{cumulate}} \]  
\[ \delta^{41} K_{\text{melt}} = \delta^{41} K_{\text{cumulate}} - \Delta^{41} K_{\text{crystal-melt}} \]

where \( f^{K}_{\text{melt}} \) is the mass fraction of K remaining in melt and \( \Delta^{41} K_{\text{crystal-melt}} \) is the K isotopic fractionation factor between crystallizing assemblage and melt. \( f^{K}_{\text{melt}} \) equals \( \frac{C_{\text{K, initial melt}}}{C_{\text{K, melt}}} \), and the relationship between the degree of crystallization (F) and K concentrations of melt (\( C_{\text{K, melt}} \)) is governed by the batch crystallization equation

\[ C_{\text{K, melt}} = \frac{C_{\text{K, initial melt}}}{(D^{K}_{\text{crystal/melt}} - 1) \times F + 1} \]

where \( D^{K}_{\text{crystal/melt}} \) and \( \Delta^{41} K_{\text{crystal-melt}} \) can be obtained by

\[ D^{K}_{\text{crystal/melt}} = \sum_{\alpha=1}^{n} (D^{\text{K, mineral } \alpha/melt} \times X_{\text{mineral } \alpha}) \]

\[ \Delta^{41} K_{\text{crystal-melt}} = \sum_{\alpha=1}^{n} (\delta^{41} K_{\text{mineral } \alpha/melt} \times X_{\text{mineral } \alpha} \times \delta^{41} K_{\text{mineral } \alpha/melt}) \]

where \( D^{\text{K, mineral } \alpha/melt} \), \( \Delta^{41} K_{\text{mineral } \alpha/melt} \), and \( X_{\text{mineral } \alpha} \) are the mineral/melt partitioning coefficient for K, mineral-melt K isotope fractionation factor, and weight portion of mineral in crystallizing assemblage, respectively.

Partitioning of K between major K-bearing minerals and peraluminous melt has been well documented (66, 67), but \( \Delta^{41} K_{\text{mineral-melt}} \) is still unknown. Considering that the K-O bond length in silicate glass (3.00 to 3.06 Å) is close to that in orthoclase (2.94 Å) (68), we used the \( \Delta^{41} K_{\text{mineral-orthoclase}} \) as an approximation to the \( \Delta^{41} K_{\text{mineral-melt}} \) in our model. The \( \delta^{41} K \) value of feldspar is inversely correlated with its orthoclase content (23, 24, 26); orthoclase-poor plagioclase (Or = 3 mol %) can be isotopically heavier than the coexisting K-rich feldspar in equilibrium (Or > 50 mol %) by 2.5‰ (24). By contrast, K isotope fractionation between mica minerals and K-rich feldspar (Or > 50%) is near zero at magmatic temperatures (23–25, 54). The feldspar in Himalayan leucogranites is composed of two nearly pure endmembers: albite-dominant plagioclase (Ab > 78 mol % and Or < 3 mol %) and orthoclase-dominant K-feldspar (Or > 83 mol %) (69). Therefore, constant \( \Delta^{41} K_{\text{plagioclase-melt}} \) of 2.5‰, \( \Delta^{41} K_{\text{K-feldspar-melt}} \) of 0‰, \( \Delta^{41} K_{\text{biotite-melt}} \) of 0‰, and \( \Delta^{41} K_{\text{muscovite-melt}} \) of 0‰ were adopted.
The modal abundance of the crystallizing mineral assemblage varies with temperature as crystallization proceeds. In the case of equilibrium crystallization, \( X_{\text{mineral}} \) is the same as the modal abundance of minerals in the average cumulate. Here, we applied the thermodynamic modeling software Rhyolite-MELTS to determine the modal mineralogy of the average cumulate throughout the crystallization (70). The calculations were performed from super-solidus (\( F = 0\%\), 900°C) to near-solidus (\( F = 90\%\), 640°C) in 0.5°C steps under the same condition as crystallization experiments on Himalayan leucogranites [pressure = 400 MPa, oxygen fugacity = -0.5 log units (71)]. Note that Rhyolite-MELTS modeling works best for metaluminous melts (70). If the phases in the Rhyolite-MELTS program were not suppressed, then our modeling produced orthopyroxene and leucite, which are never found in the Himalayan leucogranites. To better reproduce the crystallizing assemblage, only the five most common minerals (i.e., quartz, plagioclase, K-feldspar, biotite, and muscovite) in the Himalayan leucogranites were chosen as potential solid phases in our models. Other subordinate minerals such as tourmaline and garnet barely contain K, Rb, and Sr, and thus, their absence has little effect on our modeling of \( \delta^{41}K \) and Rb/Sr variations. This modeling approach predicts the earlier crystallization of plagioclase and biotite relative to K-feldspar and muscovite (Fig. 3, A to C), which is consistent with the sequence constrained by both microscopic observations and crystallization experiments on two-mica leucogranites (38, 71).

The Rb and Sr concentrations of the differentiated melt can be calculated using Eq. 5. Those of the cumulate are estimated by a mass balance equation (taking Rb as an example)

\[
C_{\text{Rb}}^{\text{cumulate}} = F \times C_{\text{Rb}}^{\text{melt}} + (1 - F) \times C_{\text{Rb}}^{\text{cumulate}}
\]

**Fractional crystallization model**

During fractional crystallization, crystals are not in equilibrium with the differentiated melt anymore once they are segregated. The modal mineralogy of the crystallizing assemblage over the differentiation sequence is estimated via the Rhyolite-MELTS modeling under the same conditions described in the equilibrium crystallization model but in the "fractionate solids" mode (70).

\[D_{\text{K/melt}}^\text{crystal/melt} \text{ and } \Delta^{41}K_{\text{crystal/melt}} \text{ are recalculated at every step of 0.5°C using Eqs. 6 and 7, and we assume a constant modal abundance of crystallizing mineral assemblage for each step. The K concentration and isotopic composition of the differentiated melt at step } i \text{ are estimated using the Rayleigh distillation equations}
\]

\[C_{\text{K/melt}}^{i} = C_{\text{K/melt}}^{i-1} \times (1 - F) \times \left( f_{\text{melt}}^{\text{crystal/melt}} \right)^{\delta^{41}K_{\text{melt}}}
\]

\[f_{\text{melt}}^{\text{crystal/melt}} = \frac{\left( M_{\text{melt}} - M_{\text{crystal/melt}} \right)}{M_{\text{melt}}} \times \frac{M_{\text{crystal/melt}}}{F} \times \frac{C_{\text{K/melt}}^{i-1}}{C_{\text{K/melt}}^{i}}
\]

\[\Delta^{41}K_{\text{melt}} = \delta^{41}K_{\text{melt}} - \delta^{41}K_{\text{crystal/melt}} \times \ln(f_{\text{melt}}^{\text{crystal/melt}})
\]

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at https://science.org/doi/10.1126/sciadv.abo4492
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