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Petrogenesis of Himalayan Leucogranites: Perspective From a Combined Elemental and Fe-Sr-Nd Isotope Study

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1. Introduction

Himalayan leucogranites are silica-rich and strongly peraluminous; they have been widely proposed to be primary melts of crustal anatexis, given their compositions comparable to experimental melts of local meta-sediments and crustal radiogenic (Sr-Nd-Pb-Hf) and stable isotopic (H-O-Si) compositions (Deniel et al., 1987; France-Lanord et al., 1988; Hopkinson et al., 2017; X.-C. Liu et al., 2018; Patiño Douce & Harris, 1998). Their petrological and geochemical variations were considered to be derived from different melting mechanisms, for example, dehydration melting or fluid-present melting of a single source (metapelite) or multiple sources (metapelite and metagraywacke or amphibolite) under different P-T-XH2O conditions (Guillot & Le Fort, 1995; Inger & Harris, 1993; Knesel & Davidson, 2002; Le Fort, 1975; Patiño Douce & Harris, 1998; Zeng, Gao, & Xie, 2011). For example, at high pressure (∼1 GPa), fluid-present melting of muscovite would generate leucogranite melts with high Na2K and CaO contents (L.-E. Gao et al., 2017; Patiño Douce & Harris, 1998; Zhang et al., 2004). However, at low pressure (∼6 kbar), muscovite dehydration melting would generate leucogranite melts with high Rb/Sr and low Na2K, Sr, and Ba contents (Harris...
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& Inger, 1992; Patiño Douce & Harris, 1998). More enriched Sr-Nd-Hf isotopes in some Himalayan intrusions suggest that the melting source contains a contribution from the Lesser Himalayan sequence (Guo & Wilson, 2012; Hopkinson et al., 2019). Some leucogranites in southern Tibet with high Sr/Y ratios and relatively depleted Sr-Nd isotopic compositions indicate partial melting of amphibolite under thickened crustal conditions (Ji et al., 2020; Z.-C. Liu et al., 2014; Zeng, Gao, & Xie, 2011; Zeng, Gao, Xie, & Liu-Zeng, 2011; Zeng et al., 2019). Thus, the Himalayan leucogranites have been used to reflect the P - T - XH2O conditions and the tectonic evolution of Himalayan orogenic belt (Aikman et al., 2008; Gao & Zeng, 2014; Hopkinson et al., 2019; Huang et al., 2017; Weinberg, 2016).

In contrast, the chemical variation of Himalayan leucogranites has been considered by some recent studies to result from late differentiation (Z.-C. Liu et al., 2014, 2019; R. Wang et al., 2017; F.-Y. Wu et al., 2020). Overall, most Himalayan leucogranites have strong negative Eu anomalies, low Zr/Hf and Nb/Ta ratios as well as a strong rare earth element tetrad effect that evolves with decreasing zircon saturation temperatures (F. Wu et al., 2017; F.-Y. Wu et al., 2020). However, these geochemical characteristics are equally consistent with varying melt fractions from sedimentary sources, an interpretation supported by petrographic evidence which precludes the fractional crystallization (FC) of feldspars in leucogranite petrogenesis (Inger & Harris, 1993). Given their highly evolved compositions and recently documented rare-metal mineralization, the leucogranites from Xiaru, Ramba, and Cuonadong plutons have been termed as highly fractionated granites (Z.-C. Liu et al., 2014, 2016; Xie et al., 2019). Accordingly, it has been proposed that most Himalayan leucogranites were generated by extreme degree of FC (e.g., four stages of 75% fractionation; F.-Y. Wu et al., 2020). As highly fractionated granites, the characteristics of primary magmas for Himalayan leucogranites are difficult to constrain (e.g., Z.-C. Liu et al., 2016). Accordingly, previous conclusions on the thermal and tectonic evolution of Himalayan orogenic belt based on leucogranites may need reconsideration. Therefore, it is crucial to elucidate whether Himalayan leucogranites were generated by extreme magma differentiation or directly by crustal anatexis. This dilemma, however, is difficult to resolve, since both petrogenesis models can explain the mineral assemblage and chemical composition of Himalayan leucogranites (e.g., Fan et al., 2021; P. Gao et al., 2016; F. Wu et al., 2017).

An iron (Fe) isotopic study could resolve this dilemma and shed new light on the petrogenesis of Himalayan leucogranites, since Fe isotopes fractionate significantly during granitic magmatism (e.g., δ56Fe from 0.08‰ to 0.64‰ for granites with SiO2 > 71 wt% in Du et al., 2017; Foden et al., 2015; He et al., 2017; Heimann et al., 2008; H. Wu et al., 2017 and Xia et al., 2017). Fe isotopic fractionation during granitic magmatism is regulated by the redox and chemical composition of the melt and co-existing mineral assemblages (Dauphas et al., 2014; He et al., 2017; Sossi & O’Neill, 2017). Phases hosting Fe with higher charge and lower coordination number prefer heavier isotopes. Mafic silicate minerals (e.g., biotite and amphibole) enriched in ferrous Fe are isotopically lighter, while magnetite (Fe3+/ΣFe = 0.69) is isotopically heavier than coexisting granitic melts with low (Na + K)/(Ca + Mg) (Heimann et al., 2008; Sossi & O’Neill, 2017; Telus et al., 2012; H. Wu et al., 2017). With increasing (Na + K)/(Ca + Mg) in the melt, its Fe isotopes can be heavier than magnetite in turn (Dauphas et al., 2014; He et al., 2017). The 56Fe of granites directly generated by crustal anatexis deviate from their source rocks by a limited extent, that is, less than Δ56Fe melt-residue, exemplified by migmatites from Black Hills and the Dabie orogen where the apparent Fe isotope fractionation between leucosomes and melanosomes is about 0.10‰ (Telus et al., 2012; L.-J. Xu et al., 2017). On the contrary, Fe isotope fractionation can become much larger during extreme FC of granitic melts, for example, elevating the melt 56Fe by > 0.5‰ (Dauphas et al., 2014; Du et al., 2017). This difference is because crustal anatexis is a batch process while FC is a Rayleigh process.

To better understand their petrogenesis, here, we report combined elemental and Fe-Sr-Nd isotopic data for 30 Himalayan leucogranites. We also measured nine metasedimentary samples from the Greater Himalayan Sequence, which represent the possible source of Himalayan leucogranites (Inger & Harris, 1993; Patiño Douce & Harris, 1998). The mineral separates from five selected leucogranite samples were also measured, because tourmaline, an important Fe hosting mineral in the leucogranite, has not been well investigated for Fe isotopes hitherto. Based on new constraints from Fe isotopic data, the role of partial melting and extreme FC has been investigated in Himalayan leucogranites.
Geological Background and Sample Description

The Himalayan orogen is a consequence of the ongoing continental collision between India and Asia since the Paleogene (Figure 1) (Ding et al., 2005; Hodges, 2000; X. Hu et al., 2016; Yin & Harrison, 2000; D. C. Zhu et al., 2015). It is divided into four lithotectonic units: (a) the Tethyan Himalayan Sequence, bounded by the Indus-Tsangpo suture to the north and the Southern Tibetan Detachment System to the south and mainly composed of very low to low-grade Neoproterozoic to Eocene marine sediments; (b) the Greater Himalayan Sequence, lying tectonically below the Tethyan Himalayan Sequence and mainly comprised of medium to high grade metasedimentary and meta-igneous rocks; (c) the Lesser Himalayan Sequence, bounded by the Main Central Thrust at the top and the Main Boundary Thrust at the bottom and made up of very-low grade to lower amphibolite facies metamorphic rocks with an age range of 1870–850 Ma; and (d) the sub-Himalayan Sequence, as the lowermost tectonic unit, mainly composed of Neogene sediments deposited in the active Himalayan foreland basin (Carosi et al., 2018; Kohn, 2014; Yin, 2006). Two roughly parallel granite belts are identified due to their temporal and spatial differences in the Himalayan orogen (Yin & Harrison, 2000). The southern one is the High Himalayan granite belt and consists of Oligocene-Miocene granite dikes, sills, and plutons intruded into the Greater Himalayan Sequence along with the Southern Tibetan Detachment System (Weinberg, 2016; F. Wu et al., 2015). The northern one is the Tethyan Himalayan granite or North Himalayan granite belt which intrudes into the Tethyan Himalayan Sequence as the North Himalayan Gneiss Dome (Zeng, Gao, Xie, & Liu-Zeng, 2011).

Thirty Himalayan leucogranites and nine metasedimentary rocks were collected in this study (Figure 1, Tables S1 and S2), and the latter represent the exposed counterparts of source rocks for the former. The leucogranite samples can be divided into three types based on their petrology: two-mica, tourmaline, and garnet leucogranites, which represent the major magmatic stage. The aplite, pegmatite, and hydrothermal alteration granite which represent the magmatic-hydrothermal and hydrothermal stages, were not considered here, given the potential complex modification on whole-rock $^{56}$Fe during hydrothermal processes (e.g., Markl et al., 2006; Y. Wang et al., 2011; B. Zhu et al., 2016). The two-mica leucogranites are composed
of biotite, muscovite, feldspars, and quartz. The other two types of leucogranites contain characteristic tourmaline and garnet instead of biotite respectively, except for sample YD1306. This sample, as a tourmaline leucogranite, also contains trace biotite. A detailed petrological description is given in Table S1. All leucogranite samples were collected distal from the pluton boundary without xenoliths and are recovered from both the Tethyan and High Himalayan granite belts (Figure 1). Metasedimentary samples were collected from the kyanite zone to the sillimanite zone of Greater Himalayan Sequence in central Nepal and Bhutan. They range from schist to gneiss with the key pelitic mineral assemblages of muscovite ± biotite ± garnet ± kyanite ± sillimanite (Table S2). The muscovite-garnet schist (Sample N13) has been used as the starting material of melting experiments that succeeded in generating melts of Himalayan leucogranite compositions (Patino Douce & Harris, 1998).

3. Analytical Methods

All the samples were thoroughly cleaned, and the weathered surface layers were removed before crushing in a corundum jaw crusher to 60 mesh. Subsequently, the samples were powdered in an agate ring mill to 200 mesh. Whole-rock major and trace elements were measured at the State Key Laboratory of Geological Processes and Mineral Resources (GPMR), Wuhan. Sample powders were mixed with LiI, LiBr, and NH₄NO₃, and fused in a Pt crucible. Major elemental data were obtained on the prepared glasses using X-ray fluorescence spectrometry (Shimadzu XRF-1800). Loss on ignition (LOI) was measured gravimetrically (Tianjin Institute of Geology and Mineral Resources following the procedure by D. Liu et al., 2015). Accuracy is better than 3% (Ma et al., 2012). For trace elements analysis, 50 mg whole-rock powders were dissolved by HF + HNO₃ in Teflon bombs, and then sealed at 190°C for 48 h. The samples were dried, dissolved in 100 g 2% HNO₃, and then measured by Agilent 7500a (Table S3). The detailed analytical procedures have been previously reported (Y. Liu et al., 2008). The accuracy is routinely better than 5% for the reported trace elements, which is confirmed by analyses of international rock standards (Table S4). The FeO contents were measured by redox titration using K₂Cr₂O₇ solution at the China University of Geosciences, Beijing, and Fe/Cr was calculated accordingly (i.e., (FeO − FeO)/FeO). The accuracy is <10% for the obtained Fe/Cr. Mineral separation was handpicked under a binocular microscope from coarsely crushed samples (80–120 mesh).

Whole-rock powders for Sr-Nd isotopes were dissolved in the clean Teflon bombs with HF + HNO₃ + HClO₄, and then chemical separations were performed using conventional ion exchange procedures. Sr and Nd isotopes were analyzed using a Thermo-Finnigan TRITON thermal ionization mass spectrometer (TIMS) at the Tianjin Institute of Geology and Mineral Resources following the procedure by D. Liu et al. (2017). Sr and Nd isotopic ratios were corrected for instrumental fractionation by normalization against ⁸⁶Sr/⁸⁸Sr = 0.1194 and ¹⁴⁶Nd/¹⁴⁴Nd = 0.7219, respectively (Table S5). Repeated analyses of Sr standards BCR-2 and NBS987 yielded average ⁸⁷Sr/⁸⁶Sr ratios of 0.704975 ± 0.000021 (2σ, n = 6) and 0.710218 ± 0.000005 (2σ, n = 7), respectively. The average ¹⁴³Nd/¹⁴⁴Nd values of BCR-2 and a laboratory internal standard solution (LRIG-Nd) are 0.512635 ± 0.000004 (2σ, n = 7) and 0.512201 ± 0.000003 (2σ, n = 8), which agrees well with the recommended values (C.-F. Li et al., 2007).

Iron isotope analyses were conducted at the Isotope Geochemistry Laboratory, China University of Geosciences, Beijing, following the procedures previously established (Dauphas, Pourmand, & Teng, 2009; He et al., 2015; C. Zhu et al., 2018). Approximately 3–35 mg whole-rock powders were dissolved in a 3:1 mixture of concentrated HF-HNO₃ in Teflon beakers on a hotplate at ~130°C until the solutions became transparent. After evaporation to dryness at 140°C, the residues were refluxed subsequently with aqua regia (HCl: HNO₃ = 3:1) and excess HNO₃ aqua regia (HCl: HNO₃ = 2:1) at 130°C for two times. The samples were dissolved in 6 N HCl before chromatographic purification. Iron was separated from matrix elements and potential isobars with 1 ml AG1-X8 pre-clean resin in an HCl medium. Whole procedure blank is less than 10 ng for Fe and thus can be considered negligible compared to >50 µg sample Fe processed. Fe isotopic ratios were analyzed on a Neptune Plus multiple collector inductively coupled plasma mass spectrometer (MC-ICP-MS) on high-resolution modes, with mass bias corrected by the sample-standard bracketing method. The Fe isotopic data are reported in δ values relative to IRMM-014 (δFe (%)) = [(Fe/⁵⁶Fe)sample/(Fe/⁵⁶Fe)IRMM-014 − 1] × 1000, where i can be 56 or 57. The isotopic measurement sequence for each sample Fe solution was repeated four times, and the mean values are reported. The internal uncertainties are given as 2SE after Dauphas, Pourmand, and Teng (2009) and He et al. (2015), considering the errors arising from both the
chemical procedures and the MC-ICP-MS measurement, which are typically ≤0.05% for δ⁵⁶Fe (Table S6). Both the long-term reproducibility and accuracy are better than 0.05% for δ⁵⁶Fe, which comes from the duplicate measurements on 24 geological standards over 7 years (He et al., 2015; C. Zhu et al., 2018). The USGS standard AGV-2 and GSP-2 were processed with unknown samples, yielding values (δ⁵⁶Fe = 0.09 ± 0.04‰ and 0.15 ± 0.04‰, respectively) consistent with the previously published values within quoted uncertainties (e.g., Craddock & Dauphas, 2011; Dauphas, Craddock, et al., 2009; He et al., 2015; Sossi et al., 2012). Duplicate analyses of samples, independent from sample dissolution to instrument analysis, also show consistent results within quoted errors (Table S6).

4. Results

The Himalayan leucogranites show large geochemical variation, especially for two-mica leucogranites which can be divided into high Sr/Y and low Sr/Y groups, based on their Sr and Y contents and Sr-Nd isotopic distinction (Figures 2a and 3). The low Sr/Y two-mica leucogranites have high SiO₂ (70.80–76.26 wt%) and Al₂O₃ (12.49–15.69 wt%) and low MgO (0.16–0.94 wt%), TiO₂ (0.09–0.33 wt%), and (Na + K)/(Ca + Mg) (2.94–15.23) values (Figure 2). They show high Rb (181–398 ppm) and varied Sr (35–262 ppm), Y (5.52–45.18 ppm), and Ba (52–615 ppm) contents, with Rb/Sr and Sr/Y ratios ranging from 0.80 to 7.32 and 0.77 to 22.39, respectively. Compared to the low Sr/Y two-mica leucogranites, the high Sr/Y two-mica leucogranites show lower Rb (84–277 ppm) and Y (2.19–8.30 ppm) and higher Sr (157–801 ppm) and Ba (202–968 ppm) contents, along with lower Rb/Sr (0.11–1.76) and higher Sr/Y (26.47–111.17) ratios (Figure 2). The tourmaline and garnet leucogranites both have TiO₂ (0.02–0.07 wt%), Sr (13–72 ppm), and Ba (6–187 ppm) contents systematically lower than the low Sr/Y two-mica leucogranite. Garnet leucogranites have the lowest MgO (14.84–27.63), Zr/Hf (11.15–19.31), and Eu/Eu* (0.10–0.29) among Himalayan leucogranites (Figure 2). The Fe³⁺/ΣFe ratios of these leucogranite samples range from 0.00 to 0.55, with an average of 0.17 and a median of 0.15 (Figure 2f), similar to that of S-type granites (~0.15) from the Lachlan Fold Belt (Chappell & White, 1992).

Sr-Nd isotope data of these Himalayan leucogranites show large variations (Table S5). The εNd(t) of low Sr/Y two-mica, tourmaline, and garnet leucogranites range from 0.7216 to 0.7674 and −16.0 to −12.4, respectively, showing affinity to the Himalayan metasedimentary rocks. However, the Sr-Nd isotopic composition of high Sr/Y leucogranite is relatively depleted (εNd(t) = 0.7088–0.7175; tNd(t) = −11.1 to −10.2) and comparable to the metabasites in THS (Figure 3 and Table S5).

Nine kyanite-zone to sillimanite-zone schists and gneisses from Bhutan and Nepal show relatively homogenous δ⁵⁶Fe ranging from 0.05‰ to 0.15‰ with an average of 0.10 ± 0.08‰ (2SD), similar to values from global clastic sediments and the mean upper continental crust (δ⁵⁶Fe ~10‰; Beard et al., 2003; Foden et al., 2015) (Table S6 and Figure 4). The tourmaline and two-mica leucogranites yield slightly isotopically heavier δ⁵⁶Fe, but with a limited range from 0.13‰ to 0.24‰ despite their large variations in petrology, elemental and Sr-Nd isotopic compositions (Figure 4). Garnet leucogranites have δ⁵⁶Fe (−0.04‰ to 0.06‰) lower than the other two types of leucogranites. Compared to the literature data, garnet leucogranites represent the lowest δ⁵⁶Fe end-member of granites at a given SiO₂.

Biotite (0.12‰–0.24‰) and tourmaline (ca. 0.15‰) have relatively consistent values among different samples, while muscovite can have variable δ⁵⁶Fe sample by sample, ranging from 0.20‰ to 0.45‰. In YD1306 where biotite co-exists with tourmaline, δ⁵⁶Fe_biotite is marginally higher than δ⁵⁶Fe_tourmaline, with a difference of 0.08 ± 0.04‰. Garnet has the lightest Fe isotopic composition with a δ⁵⁶Fe of ∼0.11‰ (Figure 5a). The similarity in δ⁵⁶Fe of the whole rock compared to biotite and tourmaline suggests that these minerals are the dominant Fe carrier in two-mica and tourmaline leucogranites respectively. This view is consistent with the observed mineral assemblages and EPMA analyses.

5. Discussion

5.1. Fe Isotopic Fractionation Factors Among Minerals and Melt

Iron isotope fractionation factors for a range of minerals and melt (Δδ⁵⁶Fe_{mineral-melt}) can be deduced from one certain mineral-melt fractionation and the inter-mineral fractionation using the equation
Δ^{56}\text{Fe}_{\text{B-melt}} = Δ^{56}\text{Fe}_{\text{A-melt}} + Δ^{56}\text{Fe}_{\text{B-A}} \quad (A \text{ and } B \text{ represent different minerals}). Δ^{56}\text{Fe}_{\text{biotite-melt}} \approx 0.1\% \) was calculated through the fractionation between the leucosomes and melanosomes of Black Hills migmatites where biotite is the dominant Fe carrier in the residua (Telus et al., 2012). This Δ^{56}\text{Fe}_{\text{biotite-melt}} can be influenced by the redox state of Fe in the melt and biotite, because phases with higher Fe^{3+}/ΣFe prefer heavier isotopes (Dauphas et al., 2014; Sossi & O’Neill, 2017). For a granitic melt, biotite tends to have similar Fe^{3+}/ΣFe to the melt under reductive conditions (Baker & Rutherford, 1996; Cesare et al., 2005). The Himalayan
leucogranites have relatively low Fe\textsuperscript{3+}/ΣFe similar to the S-type granites in the Lachlan Fold Belt (Figure 2 and Chappell & White, 1992), and also to the Black Hills migmatites inferred to have formed under reducing conditions from the presence of graphite (Nabelek, 1999). Coordination of Fe in the melt tends to be reduced leading to a larger Δ\textsuperscript{56}Fe\textsubscript{biotite-melt} with increasing SiO\textsubscript{2} and/or (Na + K)/(Ca + Mg) (Dauphas et al., 2014; Foden et al., 2015; He et al., 2017; Sossi et al., 2012). Himalayan leucogranites have comparable SiO\textsubscript{2} contents (70.80–78.36 wt%) and (Na + K)/(Ca + Mg) values (2.94–15.50) to the leucosomes of Black Hills migmatites (SiO\textsubscript{2}: 72.46–79.45 wt%; (Na + K)/(Ca + Mg): 0.08–18.37; Nabelek, 1999). Accordingly, it is reasonable to apply the Δ\textsuperscript{56}Fe\textsubscript{biotite-melt} estimated on a basis of Black Hills migmatites to the fractionation between biotite and Himalayan leucogranite melts.

The Fe isotopic fractionation factor between biotite and tourmaline has not been well-calibrated yet. Since Fe\textsuperscript{3+} occupies the octahedral site in both biotite and tourmaline (Hawthorne & Dirlam, 2011), they should have comparable Fe isotopic composition at the same Fe\textsuperscript{3+}/ΣFe ratio which is supported by the ab initio calculations (Nie et al., 2021). Tourmaline in the Himalayan leucogranite yields δ\textsuperscript{56}Fe slightly lower than biotite, with a Δ\textsuperscript{56}Fe\textsubscript{biotite-tourmaline} of 0.08 ± 0.04‰. However, the Δ\textsuperscript{56}Fe\textsubscript{biotite-tourmaline} constrained from Moosilauke metapelite is about −0.06 ± 0.03‰ at 700°C (Nie et al., 2021). This discrepancy may result from their different Fe\textsuperscript{3+}/ΣFe ratios and/or compositional variations in biotite and tourmaline (Nie et al., 2021; H. Wu et al., 2017).

Muscovite, like biotite, belongs to the mica group, and Fe in these minerals both occupies the same coordination sites (e.g., Fe\textsuperscript{3+} in the octahedral site), which suggests a Δ\textsuperscript{56}Fe\textsubscript{biotite-muscovite} near 0 if these two minerals have a similar Fe\textsuperscript{3+}/ΣFe. Δ\textsuperscript{56}Fe\textsubscript{biotite-muscovite} in the Himalayan leucogranites, however, ranges from −0.31‰ to 0.04‰ (Figure 5). This indicates either isotope disequilibrium or mineral compositional control on isotope fractionation. Fe isotope equilibrium may have not been reached between muscovite and the other minerals, possibly due to the timing of their crystallization. For example, biotite crystallizes early near the liquidus while muscovite is a near-solidus phase (Scaillet et al., 1995). Thus some muscovite may occur within Fe-free domains of quartz and feldspar representing the last formed intergranular melt (Figure S2), hence impeding Fe isotope exchange with early crystallized biotite. Muscovite may record the variably heavy Fe isotopic compositions of the intergranular melts under near solids conditions, which suggests that the δ\textsuperscript{56}Fe of melt could be significantly elevated after substantial crystallization of the leucogranites.

Given its euhedral shape with few inclusions, the garnet investigated here is magmatic (L.-E. Gao et al., 2012; Zeng et al., 2019), which is also consistent with its higher Mn contents (spessartine = 21%–29%, Grossular <1.92%; Table S7) than the xenocrystic garnets (spessartine < 5%; Harris et al., 1992) and lower Ca content than the peritectic garnet (Grossular = 11%–41%; King et al., 2011). Due to the lack of biotite in the garnet leucogranites, our data cannot provide direct constraints on the Fe isotopic fractionation between biotite and garnet. However, Δ\textsuperscript{56}Fe\textsubscript{biotite-garnet} has been previously well calibrated by measurements of biotite-garnet pairs from metamorphosed iron formations, yielding a positive Δ\textsuperscript{56}Fe\textsubscript{biotite-garnet} ~0.09 (±0.05) × 10\textsuperscript{5}°/° (Ye et al., 2020). This translates to a Δ\textsuperscript{56}Fe\textsubscript{garnet-melt} ~0.20% at 700°C, a crystallization temperature typical for Himalayan leucogranites.

Despite their low Fe contents, feldspars become important Fe carriers in high-Si granites. The Fe isotopic fractionation factors between biotite and feldspars have been calibrated from a suite of I-type granitoid in the Dabie orogen (H. Wu et al., 2017). The Δ\textsuperscript{56}Fe\textsubscript{biotite-plagioclase} and Δ\textsuperscript{56}Fe\textsubscript{biotite-alkali-feldspar} show linear relationships with the albite and orthoclase contents in plagioclase and alkali-feldspar, respectively (Δ\textsuperscript{56}Fe\textsubscript{biotite-plagioclase} = 0.015 × Ab% − 0.48; Δ\textsuperscript{56}Fe\textsubscript{biotite-alkali-feldspar} = 0.026 × Or% − 1.35), reflecting the compositional control on Fe isotope fractionation. From the typical compositions of plagioclase and K-feldspar in the Himalayan
leucogranites (Z.-C. Liu et al., 2019), the Ab% (in plagioclase) and Or% (in K-feldspar) were both assumed to be 90%, which indicates a $\Delta^{56}\text{Fe}_{\text{biotite-plagioclase}}$ of 0.77‰ and $\Delta^{56}\text{Fe}_{\text{biotite-K-feldspar}}$ of 0.89‰.

Accordingly, biotite, tourmaline, muscovite, and garnet are all predicted to be Fe isotopically lighter, while feldspars are isotopically heavier than the co-existing leucogranitic melt (summarized in Table 1).

5.2. Iron Isotope Fractionation of Himalayan Leucogranites

$\delta^{56}\text{Fe}$ of high-silica granites may be significantly elevated by the exclusion of isotopically light fluids, based on a negative correlation between $\delta^{56}\text{Fe}$ and Zr/Hf (Heimann et al., 2008; Poitrasson & Freydier, 2005). Note that melt Zr/Hf ratios may be also influenced by the dissolution/crystallization of zircon (Du et al., 2017;
L.-E. Gao et al., 2017), and thus not be applied to tracing fluid exsolution for Himalayan leucogranites. Despite this, δ56Fe does not increase with decreasing Zr/Hf from 33.59 to 11.15, nor is it correlated with either TEi,3 (the degree of the lanthanide tetrad effect) or Y/Ho (Figures 4c and 6), which together argues against a significant role of fluid exsolution in fractionating melt δ56Fe. The result here supports the previous conclusion that change in δ56Fe during fluid exsolution may not be quantitatively important for high-silica granitic melts (Du et al., 2017). Our leucogranite samples were collected far away from the pluton boundary and without xenoliths, and are spread across both the Tethyan and High Himalayan leucogranite belts, greatly exceeding the length-scale that diffusion-driven isotope fractionation may influence natural samples (Teng et al., 2011; H. Wu et al., 2018; Zambardi et al., 2014) (Figure 1). Therefore, neither chemical nor thermal diffusion will be considered here.

It has been widely documented that the Himalayan leucogranites were generated by anatexis of metasedimentary rocks, except for the high Sr/Y two-mica leucogranites which are likely to have been derived from partial melting of metabasite (Harris & Massey, 1994; Le Fort, 1981; Patiño Douce & Harris, 1998; Zeng, Gao, & Xie, 2011; Zeng, Gao, Xie, & Liu-Zeng, 2011). Recently, it has been suggested that the Himalayan leucogranites represent highly fractionated granites, derived from a high degree of FC with or without assimilation (Ji et al., 2020; Z.-C. Liu et al., 2016, 2019; Z.-Z. Wang et al., 2020; F.-Y. Wu et al., 2020). In the following section, we will discuss the petrogenesis of Himalayan leucogranites based on a combination of Fe isotopic data with elemental and Sr-Nd isotope indices.

5.2.1. Fe Isotope Evidence and a Simple Fractional Crystallization Model

In an origin model requiring a high degree of FC for Himalayan leucogranites, there are two scenarios: (a) the two-mica leucogranites represent the cumulate rocks after extracting derivative liquids which crystallized as tourmaline and garnet leucogranites (F.-Y. Wu et al., 2020); (b) the two-mica, tourmaline and garnet leucogranites represent a magma differentiation series (Ji et al., 2020; C. Liu et al., 2020; Scaillet et al., 1990; Zeng et al., 2019). Given the lighter Fe isotopic compositions at any given index of differentiation, for example, SiO2 and FeOt (Figure 4), garnet leucogranites, could not be controlled by the same mechanism as two-mica and tourmaline leucogranites and thus will be discussed separately.

Here we consider the possible effect of FC for two-mica and tourmaline leucogranites. For the Rayleigh fractionation model (Equation 1) and mass balance (Equation 2),

![Figure 5. δ56Fe of minerals separated from representative garnet, tourmaline, and two-mica leucogranites.](image-url)

### Table 1

<table>
<thead>
<tr>
<th>Mineral</th>
<th>δ56Fe in the literature</th>
<th>δ56Fe used in this study</th>
<th>Initial melt</th>
<th>Country rock</th>
</tr>
</thead>
<tbody>
<tr>
<td>Biotite</td>
<td>16.80–59.1ab</td>
<td>5.17–6.01a</td>
<td>0.028–0.099a</td>
<td>Rb (ppm)</td>
</tr>
<tr>
<td>Muskovite</td>
<td>47.4–202b</td>
<td>16–30d</td>
<td>0.013–0.121c</td>
<td>245</td>
</tr>
<tr>
<td>Magnetite</td>
<td>5.2.1.</td>
<td></td>
<td></td>
<td>280</td>
</tr>
<tr>
<td>Garnet</td>
<td>0.05</td>
<td>0.05</td>
<td>0.05</td>
<td>Sr (ppm)</td>
</tr>
<tr>
<td>Tourmaline</td>
<td>3</td>
<td>0.05</td>
<td>0.3</td>
<td>844</td>
</tr>
<tr>
<td>Plagioclase</td>
<td>4</td>
<td>4</td>
<td>6</td>
<td>40.2</td>
</tr>
<tr>
<td>K-feldspar</td>
<td>4</td>
<td>4</td>
<td>6.5</td>
<td>6.81</td>
</tr>
<tr>
<td>Initial melt</td>
<td>4</td>
<td>4</td>
<td>0.7064</td>
<td>0.8790</td>
</tr>
<tr>
<td>Country rock</td>
<td>4</td>
<td>4</td>
<td>0.14</td>
<td>0.10</td>
</tr>
</tbody>
</table>

Note. Δ56Fe_{melt} was suggested to be 0.10‰ according to the fractionation in Black Hills migmatites where Fe in the melanosome is dominated by biotite (Telus et al., 2012). Δ56Fe_{melt} and Δ56Fe_{melt} were calculated based on the Δ56Fe_{melt} in Section 5.1 and the Δ56Fe_{melt} from Ye et al. (2020). Δ56Fe_{melt} was assumed to be the same as Δ56Fe_{melt}. Δ56Fe_{melt} was calculated by the relationship of Δ56Fe_{melt} and Δ56Fe_{melt} in Section 5.2. The Sr isotopic ratios were corrected at 20 Ma. AFC, assimilation and fractional crystallization.

*Fe partition coefficients of these minerals in the literature were compiled from Icenhower and London (1995), Ewart and Griffin (1994), Sisson and Bacon (1992), Benard et al. (1985), H. Wu et al. (2017).
we can obtain:

\[
\Delta^{56}\text{Fe}_{\text{crystal-melt}} = \frac{f_{\text{Fe}} - 1}{\ln(f_{\text{Fe}})} \left( \delta^{56}\text{Fe}_{\text{average cumulate}} - \delta^{56}\text{Fe}_{\text{melt}} \right),
\]

where \( f_{\text{Fe}} \) is the residual Fe fraction in the crystallizing melt and ranges from 1.0 to 0. For the first scenario, the isotopic difference \( \Delta^{56}\text{Fe} \) between the average cumulate (\( \delta^{56}\text{Fe}_{\text{two-mica leucogranite}} = 0.17 \pm 0.06\%_\circ, \text{2SD} \)) and residual melts (\( \delta^{56}\text{Fe}_{\text{tourmaline leucogranite}} = 0.18 \pm 0.05\%_\circ, \text{2SD} \)) should be the maximum of \( \Delta^{56}\text{Fe}_{\text{crystal-melt}} \) (note that \( f_{\text{Fe}} \rightarrow 0 \)). However, this estimated \( \Delta^{56}\text{Fe}_{\text{crystal-melt}} \) is inconsistent with the mineral assemblage (dominated by mica, feldspar, and quartz; Table S1) of the assumed cumulates (i.e., two-mica leucogranites), which yields a \( \Delta^{56}\text{Fe}_{\text{crystal-melt}} \) between \(-0.05\%_\circ\) and \(-0.10\%_\circ\) (Figure 7b, see Table 1 for partition coefficients and fractionation factors which refer to discussions in Section 5.1). This scenario is therefore not applicable to the Himalayan leucogranites studied here. This view is also supported by the observation that some of the two-mica leucogranites show more evolved chemical compositions than the tourmaline leucogranites, for example, higher SiO\(_2\) and lower FeOt.

If the two-mica and tourmaline leucogranites represent a cogenetic differentiation magma series, \( \Delta^{56}\text{Fe}_{\text{crystal-melt}} \) required by a simple FC model to predict the whole rock \( \delta^{56}\text{Fe} \), can be constrained by substituting \( f_{\text{Fe}} = F_{\text{melt}} \times C_{\text{melt}} / C_{\text{melt0}} \) into the Rayleigh fractionation Equation 1, where \( F_{\text{melt}} \) is the mass fraction of residual melt. Given the difficulty of constraining \( F_{\text{melt}} \) for evolved granitic magmas, here we relate \( \delta^{56}\text{Fe}_{\text{melt}} \) to \( C_{\text{melt}} / C_{\text{melt0}} \) through:

\[
\frac{C_{\text{melt}}}{C_{\text{melt0}}} = F_{\text{melt}}^{(D-1)}.
\]

where \( D \) is the bulk partition coefficient of the crystallizing assemblage. Then we obtain:

\[
\delta^{56}\text{Fe}_{\text{melt}} = \delta^{56}\text{Fe}_{\text{melt0}} + \Delta^{56}\text{Fe}_{\text{crystal-melt}} \times \frac{D}{D - 1} \times \ln\left( \frac{C_{\text{melt}}}{C_{\text{melt0}}} \right) \left( D \neq 1 \right),
\]

Note that in the case \( D = 1 \), \( \delta^{56}\text{Fe}_{\text{melt}} \) will change with no variation in \( C_{\text{melt}} \) (i.e., constant as \( C_{\text{melt0}} \)). A linear regression between \( \delta^{56}\text{Fe}_{\text{melt}} \) and \( \ln(C_{\text{melt}} / C_{\text{melt0}}) \) indicates a slope (i.e., \( \Delta^{56}\text{Fe}_{\text{crystal-melt}} \times (D / (D - 1)) \)) of \(-0.02\) (Figure 7a). As FeOt of two-mica and tourmaline leucogranites decreases with increasing SiO\(_2\), \( D \) should
Higher than unity, which suggests a $\Delta^{56}\text{Fe}_{\text{crystal-melt}}$ between 0‰ and −0.02‰ (Figure 7b). Again, this estimate cannot be easily explained by the mineral assemblages observed in our samples (see discussions above).

It is possible that the crystallizing assemblage during magma differentiation differed from the assemblage observed in the granites. Hereafter we further test the possibility of a high degree of FC considering varying crystallizing mineral assemblages. Potential major liquidus phases include plagioclase, K-feldspar, quartz, biotite, muscovite, tourmaline, and garnet (Huang et al., 2017; Inger & Harris, 1993; Z.-C. Liu et al., 2019; Scaillet et al., 1990, 1995; Z.-Z. Wang et al., 2020). Quartz is free of Fe and thus not considered here. Biotite is considered as the only Fe-rich mineral to provide the minimum estimate of potential isotope fractionation, for the reasons below: (a) the fractionation factors of other Fe-rich minerals (i.e., muscovite, tourmaline, and garnet) are either similar to or greater than $\Delta^{56}\text{Fe}_{\text{biotite-melt}}$ (Sossi & O’Neill, 2017; Ye et al., 2020); and (b) magnetite which is isotopically heavier (Heimann et al., 2008; Sossi & O’Neill, 2017; Telus et al., 2012; H. Wu et al., 2017; Ye et al., 2020), is absent based on petrological observations and experimental results (Scaillet et al., 1995; F.-Y. Wu et al., 2020). Magnetite has neither been observed in thin sections of our samples nor has been separated by magnet from whole-rock powders. Moreover, significant crystallization of magnetite will result in a decrease in Fe/Mn along with $\delta^{56}\text{Fe}$ (Sossi et al., 2012), which is inconsistent with the observation (Figure 9b). FC from an initial melt represented by sample LZH1130 with the maximum FeOt in this study is illustrated in Figures 7c and 7d, given a crystallizing assemblage of biotite, plagioclase, and K-feldspar with variable proportions and parameters listed in Table 1. Rb/Sr of the differentiating melt is dominantly controlled by mica and feldspars and thus also considered here (Harris &

Figure 7. (a) A linear fit between the $\delta^{56}\text{Fe}$ and ln(FeOt/FeOt(i)) for two-mica and tourmaline leucogranites. (b) Diagram of estimated $\Delta^{56}\text{Fe}_{\text{mineral-melt}}$ versus $D(\text{Fe})_{\text{mineral-melt}}$. The possible Fe isotope fractionation factor between a crystallizing assemblage of quartz + plagioclase + K-feldspar + muscovite + biotite and the melt is given as the gray field. Isotopic effect of fractional crystallization is illustrated along with changes in FeOt and Rb/Sr in (c) and (d), respectively. Green-gray contour lines contour the evolved compositions with given $F_{\text{melt}}$. 

be higher than unity, which suggests a $\Delta^{56}\text{Fe}_{\text{crystal-melt}}$ between 0‰ and −0.02‰ (Figure 7b). Again, this estimate cannot be easily explained by the mineral assemblages observed in our samples (see discussions above).

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Inger, 1992). The calculated curves lead to two striking inferences: (a) the evolved melt tends to have $\delta^{56}$Fe higher than Himalayan leucogranite at a given FeOt, especially when $F_{\text{melt}}$ is low; (b) the limited $\delta^{56}$Fe variation of Himalayan leucogranite indicates a $\Delta^{56}$Fe$_{\text{crystal-melt}} \sim 0$, which implies a crystallizing assemblage of a low biotite/(biotite + feldspars) mode ratio. Extreme crystallization of such an assemblage would produce high Rb/Sr ratios that have not been observed. Overall, an extreme degree of FC, on its own can be ruled out as the origin of tourmaline and two-mica leucogranites.

5.2.2. Assimilation and Fractional Crystallization Model

Simultaneous assimilation of metasedimentary rocks with $\delta^{56}$Fe around 0.10‰ may have counteracted Fe isotope fractionation caused by FC. An assimilation and fractional crystallization (AFC) model has been previously proposed to explain the large variation of Sr-Nd isotopic ratios of Himalayan leucogranites (Ji et al., 2020). Ji et al. (2020) considered high Sr/Y samples with depleted Sr-Nd isotope compositions as the primary melt that was generated by partial melting of amphibolite, and suggested that the other leucogranites can be produced by an AFC process. Fe isotopic systematics during AFC has been illustrated in Figure 8 (DePaolo, 1981). As discussed in Section 5.2.1, a biotite + felspars crystallizing assemblage is adopted with partitioning and fractionation parameters from Table 1. $\delta^{56}$Fe of the initial melt and the country rock are assumed to be 0.14‰ and 0.10‰, typical of I-type granites (Foden et al., 2015) and Himalayan metasedimentary rocks (this study) respectively. Rb and Sr contents, as well as Sr isotopic systematics, are also considered, and elemental contents and $^{87}$Sr/$^{86}$Sr(i) of end-members are from Ji et al. (2020). Given the high FeOt of local country rocks (e.g., 6.81 wt% in Ji et al., 2020 and 4.68 ± 1.18 wt% of metasedimentary rocks here), a higher $r$ (i.e., the ratio of assimilation rate to fractional crystallization rate) requires more biotite in the crystallizing assemblage to cause a decrease in FeOt, which means a larger value for $\Delta^{56}$Fe$_{\text{crystal-melt}}$ and a higher $\delta^{56}$Fe of the evolved melt (Figure 8). The $\delta^{56}$Fe-FeOt trend of Himalayan leucogranites can only be reproduced by AFC when $r$ and $F_{\text{melt}}$ are both low, for example, $r = 0.2$ and $F_{\text{melt}} < 0.4$. In this case, the calculated melt should have an Rb/Sr substantially higher than the assumed initial melt and country rock (e.g., >100) and $^{87}$Sr/$^{86}$Sr(i) close to that of the country rock, which, however, are not consistent with observations on Himalayan leucogranites. Accordingly, a high degree of FC with simultaneous assimilation is also unable to account for the origin of tourmaline and two-mica leucogranites.

5.2.3. The Role of Crustal Anatexis

Although some degree of FC cannot be completely ruled out in the origin of Himalayan tourmaline and two-mica leucogranites, as discussed above, an extreme degree of FC is not supported by the Fe isotopic data presented in this study. $\delta^{56}$Fe and chemical compositions of these granites are unlikely to have been significantly affected after extraction from their sources. The highly variable elemental and Sr-Nd-Hf-O isotopic compositions of Himalayan leucogranites thus can be attributed to source heterogeneity and diverse melt generation reactions (Guillot & Le Fort, 1995; Inger & Harris, 1993; Knesel & Davidson, 2002; Le Fort, 1975; Patiño Douce & Harris, 1998; Zeng, Gao, & Xie, 2011). The supposed sources may range from metabasite to metasedimentary rocks for low Sr/Y two-mica and tourmaline leucogranites (Harrison, Grove, McKeegan, et al., 1999; Inger & Harris, 1993; Le Fort et al., 1987). Since metabasite and metasediments both have $\delta^{56}$Fe ~ 0.10‰ (e.g., Teng et al., 2013 and this study), $\delta^{56}$Fe of two-mica and tourmaline leucogranites are higher than their supposed sources by 0.07 ± 0.06 ‰ (2SD, N = 27). This value is consistent with isotope fractionation that was observed in migmatites from Black Hill and the Dabie orogen, that is, $\Delta^{56}$Fe$_{\text{leucosome-melanosome}}$ from 0.04‰ to 0.20‰ (Telus et al., 2012; L.-J. Xu et al., 2017), and most likely reflects isotope fractionation during crustal anatexis.

5.2.4. Origin of Low $\delta^{56}$Fe Garnet Leucogranites

Given all major Fe-rich minerals in the magmatic system, for example, biotite, muscovite, tourmaline, and garnet have $\Delta^{56}$Fe$_{\text{crystal-melt}} < 0$, the low $\delta^{56}$Fe of garnet leucogranites cannot be explained by isotope fractionation during either FC or crustal anatexis. Garnet in this type of leucogranite show euhedral shape and few inclusions with higher Mn and lower Ca contents (spessartine = 21%-29%, Grossular <1.92%; Table S7) than examples of xenocrystic garnet (spessartine < 5%; Harris et al., 1992) or peritectic garnet (Grossular = 11%-41%; King et al., 2011), thus indicating a probable magmatic origin (L.-E. Gao et al., 2012;
**Figure 8.** Illustration of an AFC process with $r = 0.2$ and 0.8. The details of primitive melt and country rock were listed in Table 1 and discussed in the text. Same as in Figure 7, Green-gray contour lines contour the evolved compositions with given $F_{\text{melt}}$. 
Zeng et al., 2019). Accumulation of this mineral in garnet leucogranites is indicated by their low (Gd/Yb)\textsubscript{N} (0.29–0.77), Fe/Mn (1.80–8.26, in mole) (Figure 9), and HREE contents higher than the other leucogranites (Figure S1). The low $\delta^{56}$Fe of garnet leucogranites thus can be explained by accumulation of magmatic garnet that hosts the lightest iron isotopes of the rock-forming minerals (Figures 5a and 7b). A simple binary mixing calculation suggests that the addition of only 2.2 wt% garnets in equilibrium with a low FeOt melt, represented by sample DL1703, can model the $\delta^{56}$Fe of garnet leucogranites down to $-0.04\%_{o}$, which is consistent with the garnet content estimation by microscope observation. Nevertheless, the mechanism for garnet crystallization remains largely unknown, possibly due to the special pristine melt composition of garnet leucogranites.

6. Conclusion

To elucidate the role of partial melting and FC in generating Himalayan leucogranites, we report Fe isotope data of 30 Himalayan leucogranites and 9 metasedimentary rocks, supplemented with mineral-pair measurement as well as geochemical and Sr-Nd isotopes data. Local metasedimentary rocks yield $\delta^{56}$Fe $\sim$ 0.10%, typical of global clastic sediments. Garnet leucogranites are isotopically lighter with $\delta^{56}$Fe ranging from $-0.04\%_{o}$ to 0.06%, possibly due to garnet accumulation. The tourmaline and two-mica leucogranites show roughly homogeneous $\delta^{56}$Fe from 0.13% to 0.24% without resolvable change with variable SiO\textsubscript{2} (70.80–78.36 wt%), MgO (0.04–0.94 wt%), FeOt (0.13–2.09 wt%), Mg\textsuperscript{#} (18.41–56.53), and Eu*/Eu (0.21–1.31). FC and AFC modeling suggest that they could not have experienced a high degree of FC. The higher $\delta^{56}$Fe relative to the supposed sources (by $\sim 0.07\%_{o}$), probably reflects isotope fractionation during crustal anatexis. This study therefore confirms that the geochemical and isotopic characteristics of Himalayan leucogranite can be used to reflect the P - T - X\textsubscript{H2O} conditions of partial melting, as indicators of the thermal and tectonic evolution of Himalayan crust and orogen.

Data Availability Statement

The supporting information involved in this study is available from the Figshare Repository (https://doi.org/10.6084/m9.figshare.14828346).

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References


Figure 9. $\delta^{56}$Fe versus (Gd/Yb)\textsubscript{N} (a) and $\delta^{56}$Fe versus Fe/Mn (in mole) (b) diagrams where the garnet leucogranites have the relatively lower $\delta^{56}$Fe, (Gd/Yb)\textsubscript{N}, and Fe/Mn values.


Heimann, A., Beard, B. L., & Johnson, C. M. (2008). The role of volatile exsolution and sub-solidus fluid/rock interactions in produc


References From The Supporting Information

